

## **Collection of cell cultures of vertebrates**

Catalogue was prepared by:

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## Species index

SPECIES	ORGAN or TISSUE	NAME OF CELL LINE
<hr/>		
<u>Cattle</u> <i>Bos taurus</i>	Kidney Trachea, embryo	MDBK (NBL-1) FBT
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<u>Chicken</u> <i>Gallus gallus</i>	Lymphoblastoma	MDCC-MSB1
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<u>Dog</u> <i>Canis familiaris</i>	Kidney	MDCK (NBL-2)
<hr/>		
<u>hamster Chinese</u> <i>Cricetulus griseus</i>	Fibrosarcoma  Lung  Ovary	B14-150  A-238 V-79 CHO-K1 DXB-11
<u>hamster Syrian</u> <i>Messocricetus Auratus</i>	Kidney	BHK-21 clone 13 HaK
<hr/>		
<u>Human</u> <i>Homo sapiens</i>	Bladder carcinoma Breast carcinoma  Burkitt lymphoma  Cervical carcinoma  Colon adenocarcinoma Colon, carcinoma Duodenum, adenocarcinoma Embryonic stem cells Epidermoid carcinoma Fibroblasts from xeroderma pigmentosum patients, SV40 virus-ttransformed Fibrosarcoma Glioblastoma Kidney hypernephroma Kidney, carcinoma Kidney, embryo Leukemia B-lymphoblastic Leukemia myelogenous  Leukemia promyelocytic Leukemia T-lymphoblastic	T-24 BT-474 Hs 578 T NAMALVA Raji Hela S 3 Hela TK <sup>-</sup> M-Hela clone 11 Caco-2 COLO 320 HSR HuTu 80 SC5 A 431  XPA HT-1080 T 98G HN OKP-GS 293 CCRF-SB KG-1 K-562 THP-1 HL-60 MOLT-3

Leukocytes	MOLT-4 Jurkat RPMI 1788
Liver adenocarcinoma	SK-HEP-1
Liver carcinoma	Hep G2
Lung carcinoma	A 549
Lung, embryo, SV40 transformed	WI-38VA13subline2RA
Lymphoma, histiocytic	U-937
Mammary gland carcinoma	BT-20 ZR-75-1 MCF-7
<u>Mesenchymal stem cells:</u>	
embryonic stem cells	SC5-MSC SC7-MSC
endometrial cells	ECL 2455 ECL 2534 ECL 2555
muscle of a limb of the embryo	M-FetMSC
the bone marrow of the embryo	FetMSC
the eyelid's skin of an adult donor	DF-1 DF-2 DF-3
the foreskin of a child	FRSN FRSN-1
pulp of a deciduous tooth	MSC-DP MSC-DP-1 MSC-DP-2
placenta	MSC-PL 2
wharton jelly of the umbilical cord	MSCWJ-1 MSCWJ-3
Myeloma	IM-9 RPMI 8226
Nasal septum carcinoma	RPMI 2650
Neuroblastoma	IMR-32 SK-N-MC
Osteosarcoma	MG-63 U-2 OS Hos (TE85, clone F5)
Osteosarcoma, chemically transformed	MNNG-HOS (TE 85, clon F-5)
Ovarian teratocarcinoma	PA-1
Pancreatic adenocarcinoma	Capan-2 AsPC-1
Pancreatic carcinoma	MIA PaCa-2 PANC-1
Rhabdomyosarcoma embryonic	RD
Rectum adenocarcinoma	SW 837
Tracheal epithelium transfected with pSVori- plasmid	CFTE 290 <sup>c</sup>
Uterine leiomyosarcoma	SK-UT-1B

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<u>Mink</u> <i>Mustela vison</i>	Lung	Mv 1 Lu (NBL-7)
<u>Monkey</u> African green <i>Cercopithecus</i> <i>aethiops</i>	Kidney	BGM CV-1 Vero Vero 76
Macaque rhesus <i>Macaca mulatta</i>	Kidney	LLC-MK2, derivative
<u>Mouse</u> <i>Mus musculus</i>	Brain, tumor Connective tissue  Fibroblasts Fibroblasts, embryo  Fibroblasts, embryo, SV40 transformed  Fibrosarcoma Glioblastoma Hepatoma  Leukemia lymphocytic Leukemia myelomonocytic lymphoid neoplasm Lymphoma  Mastocytoma Melanoma Muscle Myeloma  Neuroblastoma  Rhabdomyosarcoma	BC3H1 A-9 L-M (TK <sup>-</sup> , APRT <sup>-</sup> ) LS LSM NCTC clone 929 McCoy B 3T3 Swiss albino 3T3-Swiss J2 3T6 Swiss albino 3T3 NIH TK <sup>-</sup> BALB/3T3 clone A31 C3H10T1/2 clone 8 NIH/3T3 PA 317 Psi 2 BAG $\alpha$ STO 3T3B-SV40 3T3-SV 40 Wehi 164 EPNT-5 BWTG 3 MH-22a L 1210 Wehi-3 P388 D <sub>1</sub> EL-4 YAC-1 P-815 Clone M-3 C2C12 NSO/1 P3/NS1/1-Ag4-1(NS-1) P3X63Ag8.653 Sp2/0-Ag14 NB41A3 Neuro-2a A-7

	Sarcoma histiocytic Teratocarcinoma Testicular teratocarcinoma	MCH-7 MCH-82 J-774 P19 F9
<u>muntjac</u> <i>Muntiacus</i> <i>muntjak</i>	Skin	Indian Muntjac (M) Indian Muntjac (MT)
<u>Pig</u> <i>Sus scrofa</i>	Kidney Kidney, embryo	PK(15) SPEV
<u>Rabbit</u> <i>Oryctolagus</i> <i>Cuniculus</i>	Cornea Kidney	SIRC RK13
<u>rat kangaroo</u> <i>Potorous</i> <i>tridactylus</i>	Kidney	Pt K1 (NBL-3-11) PTK1 (NBL-3-17)
<u>Rat</u> <i>Rattus</i> <i>norvegicus</i>	Fibroblasts Ad5-transformed, embryo Fibroblasts spontaneously transformed Glioma  Hepatoma Kidney Leukemic basophilic granulocyte Leukemia basophilic chemically induced, peripheral blood Lymphosarcoma Muscle  Pancreas, insulinoma Pituitary tumor Carcoma	DFK3 K-22 2211 35 C6 HTC NRK-49F RBL-1 RBL-2H3  RLC L6J1 L-8 RIN m 5F GH3 JF 1 XCp

## Abbreviations

Ad - adenovirus  
AK - adenylate kinase  
AKTG - adrenocorticotrophin  
ATCC - American Type Culture Collection  
ATP - adenosine 5'-triphosphate  
bFGF - basic fibroblast growth factor  
BS - bovine serum  
BUdR - bromodeoxyuridine  
BVD - bovine virus diarrhea  
CSA - colony-stimulating activity  
DMEM - Dulbecco's modified Eagle's medium  
DMSO - dimethyl sulfoxide  
DNA - deoxyribonucleic acid  
DSM - German Collection of Microorganisms and Cell Cultures  
EA - early antigen  
EBNA - Epstein-Barr nuclear antigen  
EBV - Epstein-Barr virus  
ECACC - European Collection of Animal cell cultures  
ECHO - enteric cytopathogenic human orphans  
EDTA - disodium ethylene-diaminetetraacetate  
EGF - epidermal growth factor  
EMEM - minimal essential medium Eagle  
ES D - esterase - D  
ESCC - Ekaterinburg collection of continuous somatic cells of vertebrates  
FBS - fetal bovine serum  
FGF - fibroblast's growth factor  
G6PD - glucose-6-phosphate dehydrogenase  
GLO - glyoxylase  
GPRT(-) - guanine phosphoribosile transferase (-)  
HIV - Human immuno deficiency virus  
HLA - Human leucocyte antigen  
HS - horse serum  
HSV - herpes simplex virus  
HTLV - human T-cell leukemia virus  
IBR - infectious bovine rhynotracheitis  
ICLC - Interlab cell line collection  
Ig - immunoglobulin  
IL - interleukin  
LDH - lactate dehydrogenase  
Me - malic enzyme  
MNNG - methyl - N - nitroso-guanidine  
MWIEV - Russian research Inst. of Experimental veterinary  
MWIIV - D.I. Ivanovsky Institute of viriology  
NEAA - non-essential amino acids  
NK - naturally killer  
NPP - norepinephrine  
PEP - peptidase  
PGD - phosphogluconate dehydrogenase  
PGM - phosphoglucomutase  
PHA - phytohemagglutinin  
PTH - parathyroid hormone

RNA - ribonucleic acid  
SPBIC - St.Peterburg Institute of Cytology  
SPBII - St.Peterburg Institute of Influenza  
STR - short tandem repeats  
SV - simian virus  
TK - timidine kinase

## HUMAN CELL LINES

### 293 (HEK-293)

**Origin:** human, embryonal kidney , cell transformed with human adenovirus type 5 (Ad 5) DNA.

Gen. Virology 1977. 36:59; Virology 1977. 77: 319; Atlas of chromosomes of human and animal cell lines, S.E. Mamaeva, 2002. Moscow, Scientific world.

**Morphology:** epithelial-like

**Mode of cultivation:** monolayer

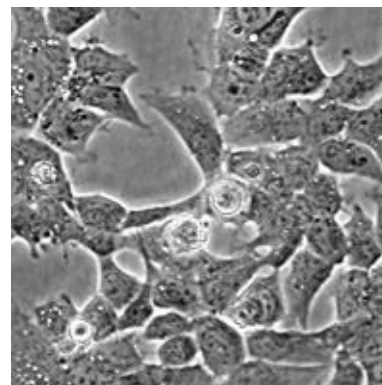
**Conditions for cultivation:** medium - EMEM

serum - FBS 10%

other components -NEAA 1%.

subculture procedure - cells detach from flask using trypsin 0.25%: EDTA 0.02% (1:2 - 1:3), split ratio 1:2 - 1:3, optimal population density  $3.0-5.0 \times 10^4$  cells/cm<sup>2</sup>, cell detach at room temperature and may take several days to reattach.

cryoconservation - growth medium, 10% DMSO,  $1.0 \times 10^6$  cells/ml in ampule



**Viability after cryoconservation:** 90-95% (0 passage, dye trypan blue)

**Sterility:** tests for bacteria, fungi and mycoplasma were negative

**Species:** karyological analysis

**Karyology:**  $2n=46$ , modal number of chromosomes 72, number of markers - 12 (differential dye), number of polyploid cells 2.4%.

**DNA profile (STR):**

Amelogenin:	X,	X
CSF1PO:	11,	12
D13S317:	12,	12
D16S539:	9,	13
D5S818:	8,	9
D7S820:	11,	12
THO1:	7,	9.3
TPOX:	11,	11
vWA:	16,	19

**Other properties:** virus susceptibility: human adenovirus type 5, astrovirus.

Contain and express the transforming genes of Ad5.

**Applications:** biotechnology (human adenovirus titration), virology, transformation

**Collections:** ATCC CRL 1573; ECACC 85120602; MWIIV; SPBIC.



**Origin:** human, epidermoid carcinoma

J.Natl.Cancer Inst. 1973. 51: 1417-1423. Atlas of chromosomes of human and animal cell lines, S.E. Mamaeva, 2002. Moscow, Scientific world.

**Morphology:** epithelial-like

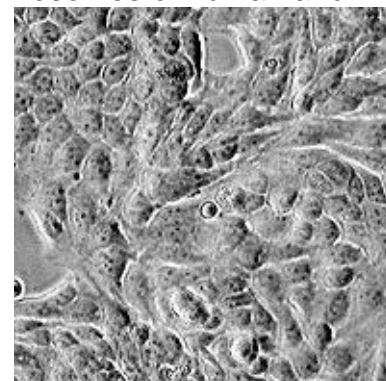
**Mode of cultivation:** monolayer

**Conditions for cultivation:** medium - DMEM

serum - FBS 10%

subculture procedure - cells detach from flask using trypsin 0.25%: EDTA 0.02% (1:3), split ratio 1:3 - 1:6, optimal population density  $2.0-4.0 \times 10^4$  cells/cm<sup>2</sup>

cryoconservation - growth medium, 10% DMSO,  $1.0 \times 10^6$  cells/ml in ampule



**Viability after cryoconservation:** 83% (0 passage, dye trypan blue)

**Sterility:** tests for bacteria, fungi and mycoplasma were negative

**Species:** karyological and isoenzymological (LDH, G6PD) analysis

**Karyology:**  $2n=46$ , variability in the range between 55-77 chromosomes, modal number of chromosomes 72, number of markers - 27 (differential dye), number of polyploid cells 7.0%

**DNA profile (STR):**

Amelogenin:	X,	X
CSF1PO:	11,	12
D13S317:	9,	13
D16S539:	12,	13, 14
D5S818:	12,	13
D7S820:	10,	10
THO1:	9,	9
TPOX:	11,	11
vWA:	15,	17

**Tumorigenicity:** tumorigenic in anti-thymocyte serum - treated NIH/Swiss mice

**Other properties:** large numbers of EGF binding sites

**Applications:** cell biology, growth factors study

**Collections:** ATCC CRL 1555; ECACC 85090402; SPBIC.

**Origin:** human, lung carcinoma.

J.Natl.Cancer Inst. 1973. 51: 1417-1423; Int.J.Cancer 1976. 17: 62-70;

Tissue Antigens 1978. 11:279.

**Morphology:** epithelial-like.

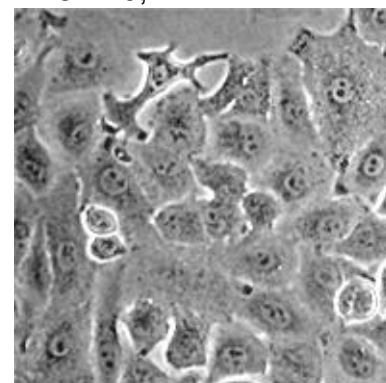
**Mode of cultivation:** monolayer.

**Conditions for cultivation:** medium - DMEM

serum - FBS 10%

subculture procedure - cells detach from flask using trypsin 0.25%: EDTA 0.02% (1:3), split ratio 1:3 -1:6, optimal population density  $2.0-4.0 \times 10^4$  cells/cm<sup>2</sup>

cryoconservation - growth medium (may add 30% BS), 5-10% DMSO,  $1.0-1.5 \times 10^6$  cells/ml in ampule.



**Viability after cryoconservation:** 97% (0 passage, dye trypan blue).

**Sterility:** tests for bacteria, fungi and mycoplasma were negative.

**Species:** karyological analysis.

**Karyology:**  $2n=46$ , variability in the range between 55-68 chromosomes, modal number of chromosomes 62-65, number of markers - 1 large submetacentric chromosome (routine dye), number of poliploid cells - 3.2%.

**DNA profile (STR):**

Amelogenin:	X,	Y
CSF1PO:	10,	12
D13S317:	11,	11
D16S539:	11,	12
D5S818:	11,	11
D7S820:	8,	11
THO1:	8,	9,3
TPOX:	8,	11
vWA:	14,	14

**Plating efficiency:** 48%.

**Tumorigenicity:** tumorigenic in nude mice.

**Other properties:** virus susceptibility: adenovirus, herpes simplex, parainfluenza II and III, polioviruses, cytomegalovirus, vesicular stomatitis.

High specific activities of choline kinase and cholinephosphate cytidyl-transferase.

Fatty acids synthesis (lecithine).

Interleukine-6 synthesis, interferon receptors.

HLA cell line phenotype F (10,w19); B (8,12).

**Applications:** biotechnology (interferon induction and titration), tumorigenicity, cell biology, enzymology, virology

**Collections:** ATCC CCL 185; ECACC 86012804; MWIIW; SPBII; SPBIC.

**Origin:** human, metastatic pancreas adenocarcinoma (ascitic fluid)

J.Natl.Cancer Inst. 1981. 67: 563-569; Clin.Lab.Med. 1982. 2: 567-578; In vitro 1982. 18: 24-34; Tumor Biol. 1985. 6: 89-98.

**Morphology:** epithelial-like

**Mode of cultivation:** monolayer

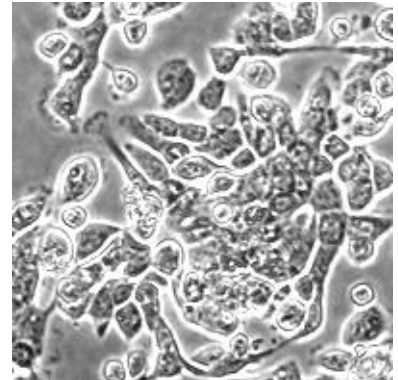
**Conditions for cultivation:** medium - RPMI 1640

serum - FBS 20%

subculture procedure - cells detach from flask using trypsin 0.25%: EDTA 0.02% (1:3), split ratio 1:2 - 1:4

cryoconservation - growth medium

10% DMSO,  $3.4 \times 10^6$  cells/ml in ampule



**Viability after cryoconservation:** 90% (0 passage, dye trypan blue)

**Sterility:** tests for bacteria, fungi and mycoplasma were negative

**Species:** karyological and isoenzymological (LDH, G6PD) analysis

**Karyology:**  $2n=46$ , modal number of chromosomes 55, number of markers – 18% cells have large submetacentric chromosome (routine dye), and 6 markers (differential dye).

**DNA profile (STR):**

Amelogenin:	X,	X
CSF1PO:	10,	13
D13S317:	9,	12
D16S539:	11,	11
D5S818:	12,	12
D7S820:	12,	13
THO1:	7,	9,3
TPOX:	8,	10
vWA:	17,	17

**Tumorigenicity:** tumorigenic in nude mice

**Applications:** tumorigenicity, immunology

**Collections:** ATCC CRL 1682; SPBIC.

**Origin:** human, mammary gland adenocarcinoma.

J. Natl. Cancer Inst. 1958. 21: 1131-1147; Int. J. Cancer 1975. 16: 74; Br. J. Cancer 2000. 83: 1309-1317; Cancer Res. 2000. 60: 4519-4525; Genes Chromosomes Cancer 2000. 28: 308-317; Atlas of chromosomes of human and animal cell lines, S.E. Mamaeva, 2002. Moscow, Scientific world.

**Morphology:** epithelial-like

Mode of cultivation: monolayer

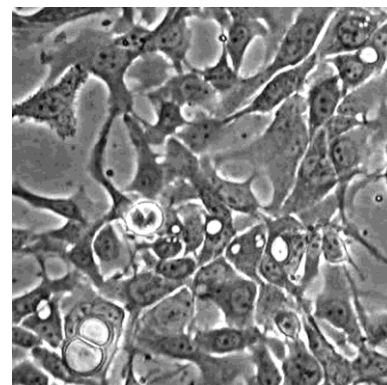
**Conditions for cultivation:** medium – EMEM

serum - FBS 10%

other components -NEAA 1%.

subculture procedure - cells detach from flask using trypsin 0.25%: EDTA 0.02% (1:3), split ratio: 1:2 -1:4

cryoconservation - growth medium, 5 - 10% DMSO,  $3.5 \times 10^6$  cells/ml in ampule



**Viability after cryoconservation:** 85% (0 passage, dye trypan blue)

**Sterility:** tests for bacteria, fungi and mycoplasma were negative.

**Species:** karyological analysis.

**Karyology:**  $2n=46$ , variability in the range between chromosomes 47-52, modal number of chromosomes 49, number of markers - 20 (differential dye), number of polyploid cells 6.5 %.

**DNA profile (STR):**

Amelogenin:	X,	X
CSF1PO:	12,	12
D13S317:	11,	11
D16S539:	11,	14
D5S818:	12,	12
D7S820:	10,	10
THO1:	7,	9.3
TPOX:	11,	11
vWA:	16,	17

**Tumorigenicity:** tumorigenic in nude mice

**Other properties:** isoenzymes: PGM3, 1; PGM1, 1; ES D, 1; AK1, 1-2; G6PD, B; GLO-1, 1-2. HLA cell phenotype A1; Bw16+/-

**Applications:** carcinogenesis, cell biology.

**Collections:** ATCC HTB 19; SPBIC.

**Origin:** human, breast, ductal carcinoma

J.Natl.Cancer Inst. 1978. 61: 967-978; In vitro 1979. 15: 723-729.

**Morphology:** epithelial-like

**Mode of cultivation:** monolayer

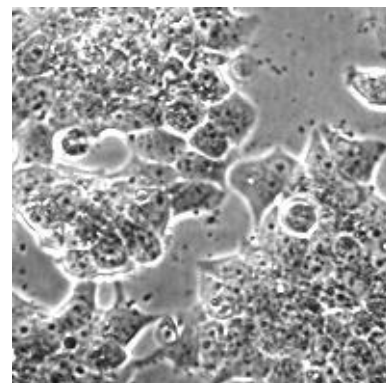
**Conditions for cultivation:** medium - RPMI 1640

serum - FBS 10%

other components - bovine insulin 10  
μ/ml

subculture procedure - cells detach from  
flask using trypsin 0.25%: EDTA 0.02%  
(1:3), split ratio 1:2-1:4

cryoconservation - growth medium 10%  
DMSO,  $5.0 \times 10^6$  cells/ml in ampule



**Viability after cryoconservation:** 71% (0 passage, dye trypan blue)

**Sterility:** tests for bacteria, fungi and mycoplasma were negative

**Species:** karyological analysis

**Karyology:**  $2n=46$ , variability in the range between 95-107 chromosomes, modal number of chromosomes 100-103, number of markers - 1 large submetacentric chromosome (routine dye), and 9 markers (differential dye), number of poliploid cells 0.2 %

**DNA profile (STR):**

Amelogenin:	X,	X
CSF1PO:	10,	11
D13S317:	11,	11
D16S539:	9,	11
D5S818:	11,	13
D7S820:	9,	12
THO1:	7,	7
TPOX:	8,	8
vWA:	15,	16

**Tumorigenicity:** tumorigenic in nude mice

**Other properties:** virus susceptibility: mouse mammary tumor virus R-III-MuMTV; isoenzymes G6PD, B; PGM<sub>1</sub>, 1; PGM<sub>3</sub>, 1; ES D, 1; Me-2, 0; AK1, 1; GLO-1, 1; R-III-MuMTV replication.

**Applications:** tumorigenicity, virology, cell biology

**Collections:** ATCC HTB 20; SPBIC.

## Caco-2

**Origin:** human, colon adenocarcinoma

J. Natl.Cancer Inst. 1977. 58: 209-214; J. Natl.Cancer Inst. 1977. 59: 221-226.

**Morphology:** epithelial-like

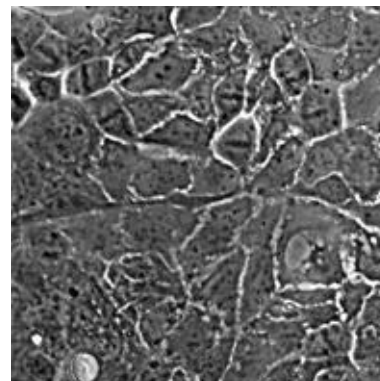
**Mode of cultivation:** monolayer

**Conditions for cultivation:** medium - DMEM

serum - FBS 10-15%

subculture procedure - cells detach from flask using trypsin 0.25%: EDTA 0.02% (1:1 - 1:3), split ratio 1:2 - 1:4, optimal population density  $2.0-4.0 \times 10^4$  cells/cm<sup>2</sup>

cryoconservation - growth medium, 5-10% DMSO,  $1.0 \times 10^6$  cells/ml in ampule



**Viability after cryoconservation:** 80% (0 passage, dye trypan blue)

**Sterility:** tests for bacteria, fungi and mycoplasma were negative

**Species:** karyological and isoenzymological (LDH, G6PD) analysis (ATCC)

**Karyology:**  $2n=46$ , variability in the range between 91-107 chromosomes, modal number of chromosomes 96-101, number of markers - 10 (differential dye), number of poliploid cells 3.2%.

**DNA profile (STR):**

Amelogenin:	X,	X
CSF1PO:	11,	11
D13S317:	11,	13, 14
D16S539:	12,	13
D5S818:	12,	13
D7S820:	11,	12
THO1:	6,	6
TPOX:	9,	11
vWA:	16,	18

**Tumorigenicity:** tumorigenic in nude mice

**Other properties:** isoenzymes Me-2,1; PGM<sub>3</sub>,1; PGM<sub>1</sub>, 1; ES D,1; AK 1,1; GLO-1,1; G6PD, B.

Lipid production.

**Applications:** gastroenterology, biochemistry, tumorigenicity, cell biology, biophysics.

**Collections:** ATCC HTB 37; ECACC 86010202; SPBIC.

## Capan-2

**Origin:** human, pancreas adenocarcinoma.

Submitted by ATCC 1990.

**Morphology:** polygonal

**Mode of cultivation:** monolayer

**Conditions for cultivation:** medium - RPMI 1640

serum - FBS 10%

subculture procedure - cells detach from flask using trypsin 0.25%: EDTA 0.02% (1:4), split ratio 1:2 - 1:4

cryoconservation - growth medium, 5-10% DMSO,  $2.0 \times 10^6$  cells/ml in ampule

**Viability after cryoconservation:** 92 % (0 passage, dye trypan blue)

**Sterility:** tests for bacteria, fungi and mycoplasma were negative

**Species:** karyological and isoenzymological (LDH, G6PD) analysis

**Karyology:**  $2n=46$ , variability in the range between 63-71, modal number of chromosomes 68-70, number of poliploid cell 2.0%.

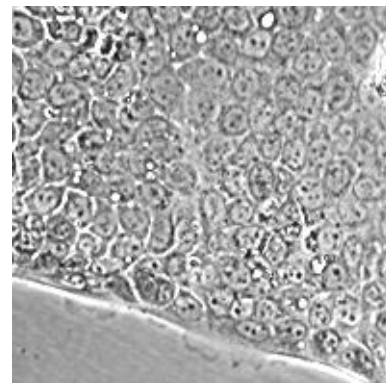
<b>DNA profile (STR):</b>	Amelogenin:	X	X
	CSF1PO:	11	12
	D13S317:	11	12
	D16S539:	9	13
	D5S818:	11	12
	D7S820:	9	11
	THO1:	9.3	9.3
	TPOX:	8	8
	vWA:	17	17

**Tumorigenicity:** tumorigenic in nude mice

**Other properties:** isoenzymes Me-2, 2; PGM<sub>3</sub>, 2; PGM<sub>1,1</sub>; ES D,1; AK1,1; GLO-1, 2; G6PD, B.

**Applications:** tumorigenicity, immunology, biochemistry.

**Collections:** ATCC HTB 80; SPBIC.



**Origin:** human, acute B-lymphoblastic leukemia, peripheral blood  
Cancer Res. 1967. 27: 2479-24-82; Atlas of

chromosomes of human and animal cell lines, S.E.

Mamaeva, 2002. Moscow, Scientific world

**Morphology:** lymphoblast-like

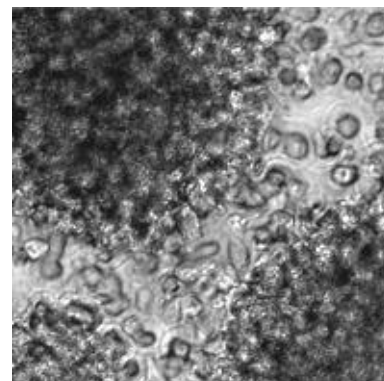
**Mode of cultivation:** suspension

**Conditions for cultivation:** medium - RPMI 1640

serum - FBS 10%

subculture procedure - optimal population  
density  $5.0 \times 10^5$  cells/cm<sup>2</sup>

cryoconservation - growth medium 5-10%  
DMSO,  $3.0-4.0 \times 10^6$  cells/ml in ampule



**Viability after cryoconservation:** 75% (0 passage, dye  
trypan blue)

**Sterility:** tests for bacteria, fungi and mycoplasma were negative

**Species:** karyological analysis

**Karyology:**  $2n=46$ , variability in the range between 42-47, modal number of  
chromosomes 46, diploid, normal human karyotype (46, XY). Number of poliploid cells  
1%.

**DNA profile (STR):**

Amelogenin:	X,	Y
CSF1PO:	10,	12
D13S317:	10,	12
D16S539:	9,	13
D5S818:	11,	12
D7S820:	11,	12
THO1:	9,	10
TPOX:	8,	8
vWA:	18,	18

**Other properties:** Ig non synthesised.

Isoenzymes - G6PD, B.

Erythrocyte rosette tests: E, 0; EA, 6%; EAC, 23%.

HLA cell line phenotype A1, A2, B12, B17, Cw2.

Positive for EBNA

**Applications:** immunology, cell biology.

**Collections:** ATCC CCL 120; ECACC 89090405; SPBIC.



**Origin:** human, tracheal epithelium, cells were transfected with pSVori- plasmid.  
Am.J.Respir.Cell Mol.Biol. 1993. 8; 522-529.

**Morphology:** epithelial-like

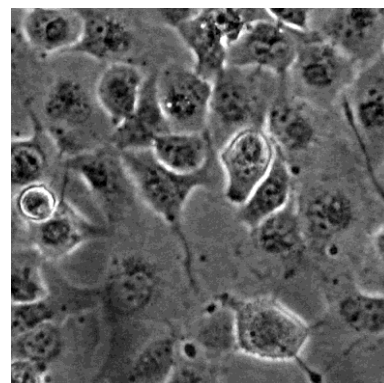
**Mode of cultivation:** monolayer

**Conditions for cultivation:** medium - DMEM

serum - FBS 10%

subculture procedure - cells detach from flask using trypsin 0.25%: EDTA 0.02% (1:3), split ratio 1:2 - 1:4

cryoconservation - growth medium, 10% DMSO,  $1.5 \times 10^6$  cells/ml in ampule



**Viability after cryoconservation:** 90% (0 passage, dye trypan blue)

**Sterility:** tests for bacteria, fungi and mycoplasma were negative

**Species:** karyological and isoenzymological (LDH, G6PD) analysis

**Karyology:**  $2n = 46$ , variability in the range between 65-73 chromosomes, modal number of chromosomes 69-70, number of markers – 24% dicentrics (routine dye); number of poliploid cells 3.5%.

**DNA profile (STR):**

Amelogenin:	X,	X
CSF1PO:	10,	13
D13S317:	9,	11
D16S539:	10,	12
D5S818:	11,	12
D7S820:	10,	11
THO1:	7,	7
TPOX:	8,	11
vWA:	17,	17

**Plating efficiency:** 30%

**Other properties:** keratin expression.

Homozygous  $\Delta$  F508-mutation (cystic fibrosis - recessive genetical disease)

**Applications:** genetical transformation and hereditary diseases studies, cell biology.

**Collections:** SPBIC.

## COLO 320 HSR

**Origin:** human, colon, carcinoma.

Cancer Res. 1979. 39: 4914; Atlas of chromosomes of human and animal cell lines, S.E. Mamaeva, 2002. Moscow, Scientific world.

**Morphology:** round cells

**Mode of cultivation:** semisuspension

**Conditions for cultivation:** medium - RPMI 1640

serum - FBS 10%

subculture procedure - split ratio 1:3,  
optimal population density  $3.0-9.0 \times 10^5$   
cells/cm<sup>2</sup>

cryoconservation - growth medium, 10%  
DMSO,  $3.0 \times 10^6$  cells/ml in ampule

**Viability after cryoconservation:** 85% (0 passage, dye trypan blue)

**Sterility:** tests for bacteria, fungi and mycoplasma were negative

**Species:** karyological and isoenzymological (LDH, G6PD) analysis

**Karyology:**  $2n=46$ , variability in the range between 49-61 chromosomes, modal number of chromosomes 52, markers - 18 (differential dye), double minute chromosomes.

**DNA profile (STR):**

Amelogenin:	X,	X
CSF1PO:	11,	11
D13S317:	11,	11
D16S539:	11,	12
D5S818:	12,	12
D7S820:	9,	12
THO1:	8,	9
TPOX:	8,	9
vWA:	15,	18

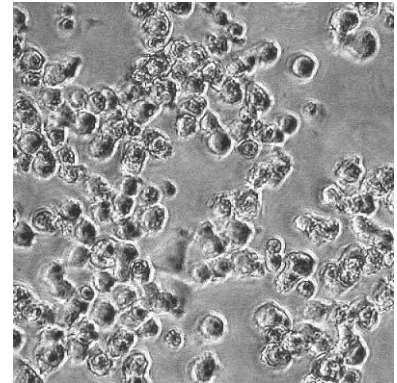
**Plating efficiency:** 12%.

**Tumorigenicity:** tumorigenic in nude mice

**Other properties:** isoenzymes PGM<sub>1,1</sub>; PGM<sub>3,1</sub>; G6PD, B; PEP-D,1; PGD, A; ES D, 1  
Serotonin, epinephrine, AKTG, NPP, PTH production

**Applications:** biochemistry, biophysics, endocrinology.

**Collections:** ATCC CCL 220.1; ECACC 87101501; SPBIC.



**Origin:** human, mesenchymal stem cells from eyelid's skin of of 37 year old woman. Tsitologiya. 2016. 57 (11): 850 – 864

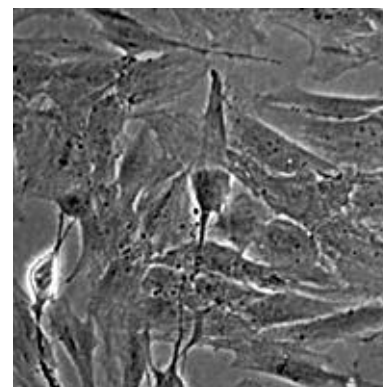
**Morphology:** fibroblast-like

**Mode of cultivation:** monolayer

**Conditions for cultivation:** medium – DMEM/F12  
serum - FBS 10%

subculture procedure - cells detach from flask using trypsin 0.25%: EDTA 0.02% (1:3), split ratio: 1:3, optimal population density 4.0- 5.0x10<sup>4</sup> cells/cm<sup>2</sup>

cryoconservation - growth medium, 10% DMSO, 1.0-1.5.x10<sup>6</sup> cells/ml in ampule



**Viability after cryoconservation:** 85% (0 passage, dye trypan blue)

**Sterility:** tests for bacteria, fungi and mycoplasma were negative.

**Species:** karyological analysis.

**Karyology:** 2n=46, modal number of chromosomes 46 (98.0%), normal human karyotype (46, XX), detected nonclonal structural chromosome rearrangements, number of poliploid cells 0.8%.

**DNA profile (STR):**

Amelogenin:	X,	X
CSF1PO:	11,	11
D13S317:	11,	11
D16S539:	10,	12
D5S818:	9,	13
D7S820:	10,	12
THO1:	9.3,	9.3
TPOX:	8,	9
vWA:	15,	19

**Plating Efficiency:** 34.5%

**Other properties:** finite lifetime culture; average population doubling time 40.0 h; The presence of surface antigens specific for human MSCs: CD44, CD73, CD90, CD105, HLA-ABC, vimentin and the lack of CD34, HLA-DR; The ability to induced differentiation into osteogenic, chondrogenic and adipogenic directions.

**Applications:** cell biology, biotechnology, feeder for cultivation embryonic stem cells.

**Collections:** SPBIC.

**Origin:** human, mesenchymal stem cells from eyelid's skin of 45 year old woman  
Tsitologiya. 2016. 57 (11): 850 – 864

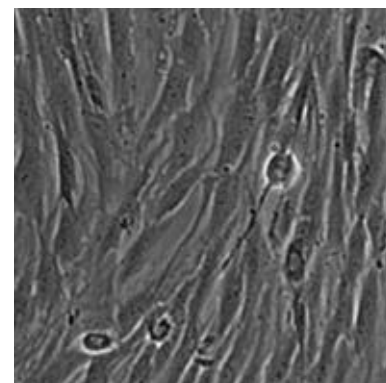
**Morphology:** fibroblast-like

**Mode of cultivation:** monolayer

**Conditions for cultivation:** medium – DMEM/F12  
serum - FBS 10%

subculture procedure - cells detach from flask using trypsin 0.25%: EDTA 0.02% (1:3), split ratio: 1:3, optimal population density 4.0- 5.0x10<sup>4</sup> cells/cm<sup>2</sup>

cryoconservation - growth medium, 10% DMSO, 1.0-1.5.x10<sup>6</sup> cells/ml in ampule



**Viability after cryoconservation:** 85% (0 passage, dye trypan blue)

**Sterility:** tests for bacteria, fungi and mycoplasma were negative.

**Species:** karyological analysis.

**Karyology:** 2n=46, modal number of chromosomes 46 (98.0%), normal human karyotype (46, XX), number of poliploid cells 1.2%.

**DNA profile (STR):**

Amelogenin:	X,	X
CSF1PO:	10,	12
D13S317:	11,	11
D16S539:	11,	11
D5S818:	11,	13
D7S820:	13,	13
THO1:	6,	9
TPOX:	9,	9
vWA:	15,	17

**Plating Efficiency:** 25.4%

**Other properties:** finite lifetime culture; average population doubling time 40.0 h; The presence of surface antigens specific for human MSCs: CD44, CD73, CD90, CD105, HLA-ABC, vimentin and the lack of CD34, HLA-DR; The ability to induced differentiation into osteogenic, chondrogenic and adipogenic directions.

**Applications:** cell biology, biotechnology, feeder for cultivation embryonic stem cells.

**Collections:** SPBIC.

**Origin:** human, mesenchymal stem cells from eyelid's skin of 53 year old woman. Tsitologiya. 2016. 57 (11): 850 – 864

**Morphology:** fibroblast-like

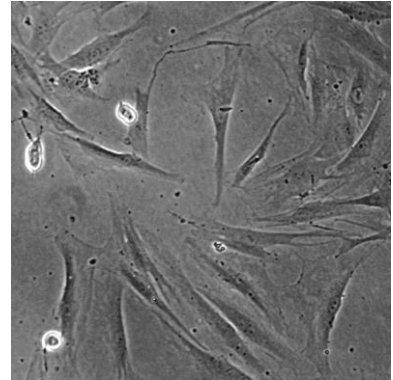
**Mode of cultivation:** monolayer

**Conditions for cultivation:** medium – DMEM/F12

serum - FBS 10%

subculture procedure - cells detach from flask using trypsin 0.25%: EDTA 0.02% (1:10), split ratio: 1:3, optimal population density 4.0- 5.0x10<sup>4</sup> cells/cm<sup>2</sup>

cryoconservation - growth medium, 10% DMSO, 1.0-1.5.x10<sup>6</sup> cells/ml in ampule



**Viability after cryoconservation:** 85% (0 passage, dye trypan blue)

**Sterility:** tests for bacteria, fungi and mycoplasma were negative.

**Species:** karyological analysis.

**Karyology:** 2n=46, modal number of chromosomes 46 (98.0%), normal human karyotype (46, XX), analysis of 3 independent replicates from one cell population revealed non-clonal structural chromosomal rearrangements (14.0%), number of poliploid cells 0.8%.

**DNA profile (STR):**

Amelogenin:	X,	X
CSF1PO:	11,	13
D13S317:	8,	12
D16S539:	11,	12
D5S818:	12,	13
D7S820:	11,	12
THO1:	6,	8
TPOX:	8,	11
vWA:	15,	16

**Plating Efficiency:** 30.0%

**Other properties:** Finite lifetime culture; average population doubling time 33.0 h; The time of the active (logarithmic) growth phase is 48 h. The stage of active replicative senescence begins at passage 25

The presence of surface antigens specific for human MSCs: CD44, CD73, CD90, CD105, HLA-ABC, vimentin and the lack of CD34, CD45, HLA-DR; The ability to induced differentiation into osteogenic, chondrogenic and adipogenic directions.

**Applications:** cell biology, biotechnology.

**Collections:** SPBIC.

**Origin:** human, endometrial cells (mesenchymal stem cells), isolated from a biopsy of a woman without endometriosis, but with a concomitant disease –endometrial polyp.

Tsitologiya. 2019. 61 (11): 902-914.

**Morphology:** fibroblast-like

**Mode of cultivation:** monolayer

**Conditions for cultivation:**

medium – IMDM or DMEM/F12 (1:1)

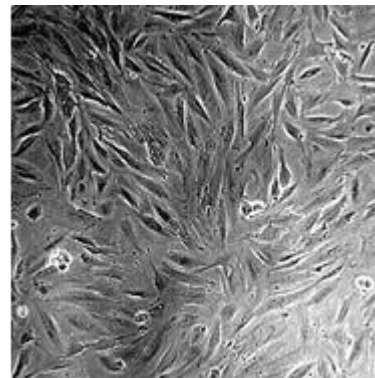
serum - FBS 10%

subculture procedure - cells detach from flask using trypsin

0.25% : EDTA 0.02%, split ratio: 1:3, optimal population

density  $4.5\text{-}5.0 \times 10^3$  cells/cm<sup>2</sup>

cryoconservation - growth medium, 10% DMSO,  $0.55\text{-}0.70 \times 10^6$  cells/ml in ampule.



**Viability after cryoconservation:** 75% (0 passage, dye trypan blue)

**Sterility:** tests for bacteria, fungi and mycoplasma were negative.

**Species:** karyological analysis.

**Karyology:**  $2n=46$ , modal number of chromosomes 46 ( $93.0 \pm 2.6$  %), normal human karyotype (46, XX), The proportion of polyploidy cells is  $21.8 \pm 4.1$ %.

**DNA profile (STR):**

Amelogenin:	X,	X
CSF1PO:	10,	12
D13S317:	11,	12
D16S539:	12,	12
D5S818:	12,	13
D7S820:	11,	11
THO1:	7	9,3
TPOX:	8	8
vWA:	14	17

**Other properties:** finite lifetime culture; The stage of active replicative senescence occurs at 18 passage.

The presence of surface antigens specific for human MSCs: CD44, CD73, CD90, CD105, HLA-ABC and the lack of CD34, CD45, HLA-DR.

Expression of estrogen and progesterone receptors (immunocytochemistry).

The ability to induced differentiation into osteogenic, adipogenic and chondrogenic directions; directional differentiation in the deductive direction was confirmed, induced by combination of hormones – estrogen (10 nM) and progesterone (1 $\mu$ M) for 14 days.

**Applications:** cell biology, biotechnology.

**Collections:** SPBIC.

**Origin:** human, endometrial cells (mesenchymal stem cells), isolated from a biopsy of a woman without endometriosis, but with a concomitant disease – uterine fibroids  
Tsitolgiya. 2019. 61 (11): 902-914.

**Morphology:** fibroblast-like

**Mode of cultivation:** monolayer

**Conditions for cultivation:**

medium – IMDM or DMEM/F12 (1:1) миома матки

serum - FBS 10%

subculture procedure - cells detach from flask using trypsin

0.25% : EDTA 0.02%, split ratio: 1:2 -1:3, optimal

population density 5.5.- 6.5x10<sup>3</sup> cells/cm<sup>2</sup>

cryoconservation - growth medium, 10% DMSO, 0.75-0.80x10<sup>6</sup> cells/ml in ampule.



**Viability after cryoconservation:** 65-70% (0 passage, dye trypan blue)

**Sterility:** tests for bacteria, fungi and mycoplasma were negative.

**Species:** karyological analysis.

**Karyology:** 2n=46, modal number of chromosomes 46 (96.0 ± 2.8 %), normal human karyotype (46, XX), The proportion of polyploidy cells is 7.0±3.6%.

**DNA profile (STR):**

Amelogenin:	X,	X
CSF1PO:	10,	11
D13S317:	11,	11
D16S539:	12,	13
D5S818:	7,	11
D7S820:	10,	11
THO1:	7	9,3
TPOX:	8	10
vWA:	16	18

**Other properties:** finite lifetime culture; The stage of active replicative senescence occurs at 17 passage.

The presence of surface antigens specific for human MSCs: CD44, CD73, CD90, CD105, HLA-ABC and the lack of CD34, CD45, HLA-DR.

Expression of estrogen and progesterone receptors (immunocytochemistry).

The ability to induced differentiation into osteogenic, adipogenic and chondrogenic directions; directional differentiation in the deductive direction was confirmed, induced by combination of hormones – estrogen (10 nM) and progesterone (1μM) for 14 days.

**Applications:** cell biology, biotechnology.

**Collections:** SPBIC.

**Origin:** human, endometrial cells (mesenchymal stem cells), isolated from a biopsy of a woman without endometriosis, but with a concomitant disease – left ventricular dermoid cyst.

Tsitologiya. 2019. 61 (11): 902-914.

**Morphology:** fibroblast-like

**Mode of cultivation:** monolayer

**Conditions for cultivation:**

medium – IMDM or DMEM/F12 (1:1)

serum - FBS 10%

subculture procedure - cells detach from flask using trypsin

0.25% : EDTA 0.02%, split ratio: 1:3, optimal population

density  $4.5 - 5.5 \times 10^3$  cells/cm<sup>2</sup>

cryoconservation - growth medium, 10% DMSO, 0.55-

$0.75 \times 10^6$  cells/ml in ampule.

**Viability after cryoconservation:** 65% (0 passage, dye trypan blue)

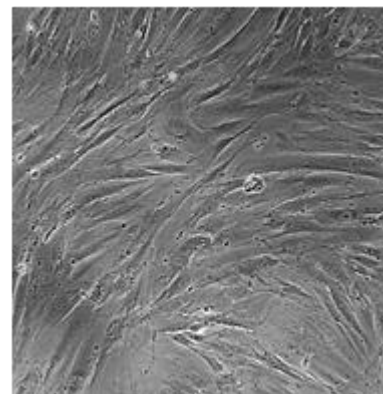
**Sterility:** tests for bacteria, fungi and mycoplasma were negative.

**Species:** karyological analysis.

**Karyology:**  $2n=46$ , modal number of chromosomes 46 ( $97.0 \pm 1.7$  %), normal human karyotype (46, XX), The proportion of polyploidy cells is  $14.0 \pm 3.5$ %.

**DNA profile (STR):**

Amelogenin:	X,	X
CSF1PO:	9,	12
D13S317:	8,	9
D16S539:	11,	12
D5S818:	12,	12
D7S820:	11,	12
THO1:	7	9
TPOX:	8	8
vWA:	17	20



**Other properties:** finite lifetime culture; The stage of active replicative senescence occurs at 18 passage.

The presence of surface antigens specific for human MSCs: CD44, CD73, CD90, CD105, HLA-ABC and the lack of CD34, CD45, HLA-DR.

Expression of estrogen and progesterone receptors.

The ability to induced differentiation into osteogenic, adipogenic and chondrogenic directions; directional differentiation in the deductive direction was confirmed, induced by combination of hormones – estrogen (10 nM) and progesterone (1 $\mu$ M) for 14 days.

**Applications:** cell biology, biotechnology.

**Collections:** SPBIC.



**Origin:** human, mesenchymal stem cells from bone marrow of 5-6 week embryo.

Tsitologiya. 2012. 54 (1): 5 – 16; Tsitologiya. 2014. 56 (8): 562 – 573.

**Morphology:** fibroblast-like

**Mode of cultivation:** monolayer

**Conditions for cultivation:** medium – DMEM/F12 serum - FBS 10%

subculture procedure - cells detach from flask using trypsin 0.25%: EDTA 0.02% (1:3), split ratio: 1:3, optimal population density 4.0-5.0x10<sup>4</sup> cells/cm<sup>2</sup>

cryoconservation - growth medium, 10% DMSO, 1.5-2.0x10<sup>6</sup> cells/ml in ampule

**Viability after cryoconservation:** 85% (0 passage, dye trypan blue)

**Sterility:** tests for bacteria, fungi and mycoplasma were negative.

**Species:** karyological analysis.

**Karyology:** 2n=46, modal number of chromosomes 46 (97.0±1.7%), normal human karyotype (46, XY), number of poliploid cells 3.0%.

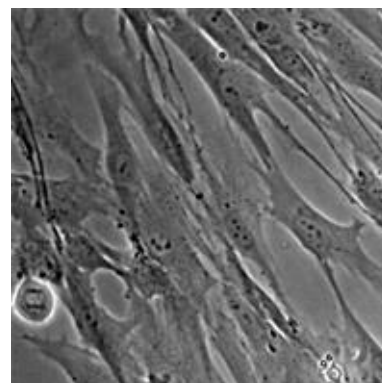
**ДНК профиль (STR):**

Amelogenin:	X,	Y
CSF1PO:	9,	12
D13S317:	11,	12
D16S539:	11,	11
D5S818:	12,	13
D7S820:	10,	12
THO1:	7,	8
TPOX:	8,	11
vWA:	14,	15

**Other properties:** finite lifetime culture; average population doubling time 33.5 h; The presence of surface antigens specific for human MSCs: CD44, CD73, CD90, CD105, HLA-ABC, vimentin and the lack of CD34, HLA-DR; The ability to induced differentiation into osteogenic, chondrogenic and adipogenic directions.

**Applications:** cell biology, biotechnology, feeder for cultivation embryonic stem cells.

**Collections:** SPBIC.



**Origin:** human, mesenchymal stem cells from foreskin of a 3-years-old boy.

Tsitologiya. 2012. 54 (1): 5 –16.

**Morphology:** fibroblast-like

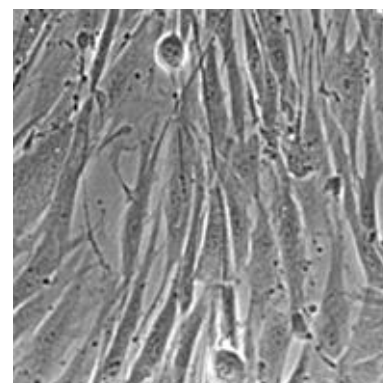
**Mode of cultivation:** monolayer

**Conditions for cultivation:** medium – IMDM

serum - FBS 10%

subculture procedure - cells detach from flask using trypsin 0.25%: EDTA 0.02% (1:3), split ratio: 1:3, optimal population density 2.0- 4.0x10<sup>4</sup> cells/cm<sup>2</sup>

cryoconservation - growth medium, 5% DMSO, 1.5-2.0x10<sup>6</sup> cells/ml in ampule



**Viability after cryoconservation:** 90% (0 passage, dye trypan blue)

**Sterility:** tests for bacteria, fungi and mycoplasma were negative.

**Species:** karyological analysis.

**Karyology:** 2n=46, modal number of chromosomes 46 (98.5±1.2%), normal human karyotype (46, XY), number of poliploid cells 13.0%.

**DNA profile (STR):**

Amelogenin:	X,	Y
CSF1PO:	10,	10
D13S317:	8,	11, 12
D16S539:	12,	13, 14
D5S818:	12,	12
D7S820:	8,	9, 12
THO1:	6,	6
TPOX:	8,	8
vWA:	16,	17, 18

**Other properties:** finite lifetime culture; average population doubling time 30.0 h; The presence of surface antigens specific for human MSCs: CD44, CD73, CD90, CD105, HLA-ABC and the lack of CD34, HLA-DR; The ability to induced differentiation into osteogenic, chondrogenic and adipogenic directions.

**Applications:** cell biology, biotechnology, feeder for cultivation embryonic stem cells.

**Collections:** SPBIC.

**Origin:** human, mesenchymal stem cells from foreskin of a 2.5 -years-old boy.  
Tsitologiya. 2016. 60 (4): 262 – 272.

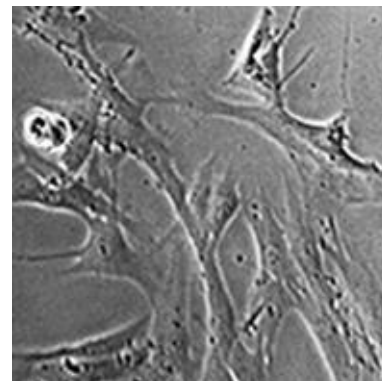
**Morphology:** fibroblast-like

**Mode of cultivation:** monolayer

**Conditions for cultivation:** medium – IMDM  
serum - FBS 10%

subculture procedure - cells detach from flask using trypsin 0.25%: EDTA 0.02% (1:3), split ratio: 1:3-1:4, optimal population density 2.0- 4.0x10<sup>4</sup> cells/cm<sup>2</sup>

cryoconservation - growth medium, 5% DMSO, 1.0-1.5x10<sup>6</sup> cells/ml in ampule



**Viability after cryoconservation:** 90% (0 passage, dye trypan blue)

**Sterility:** tests for bacteria, fungi and mycoplasma were negative.

**Species:** karyological analysis.

**Karyology:** 2n=46, modal number of chromosomes 46 (98.0±1.4%), normal human karyotype (46, XY), number of poliploid cells 6.4 %.

**DNA profile (STR):**

Amelogenin:	X,	Y
CSF1PO:	11,	11
D13S317:	9,	11
D16S539:	11,	11
D5S818:	12,	13
D7S820:	10,	12
THO1:	9,	9.3
TPOX:	11,	11
vWA:	13,	16

**Plating efficiency:** 25.1 %

**Other properties:** finite lifetime culture; average population doubling time 36.9 h. The presence of surface antigens specific for human MSCs: CD44, CD73, CD90, CD105, HLA-ABC and the lack of CD34, HLA-DR; The ability to induced differentiation into osteogenic, chondrogenic and adipogenic directions.

**Applications:** cell biology, biotechnology.

**Collections:** SPBIC.

**Origin:** human, epithelioid cervical carcinoma, strain of HeLa  
Atlas of chromosomes of human and animal cell lines, S.E. Mamaeva, 2002. Moscow, Scientific world.

**Morphology:** round and epithelial-like

**Mode of cultivation:** semisuspension

**Conditions for cultivation:** medium - EMEM

serum - FBS10%

other components - NEAA 1%

subculture procedure - optimal population density 3.0-9.0x10<sup>5</sup> cells/ml

cryoconservation - growth medium, 5%DMSO, 3.0x10<sup>6</sup> cells/ml in ampule

**Viability after cryoconservation:** 90% (0 passage, dye trypan blue)

**Sterility:** tests for bacteria, fungi and mycoplasma were negative

**Species:** karyological and isoenzymological (LDH, G6PD) analysis

**Karyology:** 2n=46, variability in the range between 51-74 chromosomes modal number of chromosomes 66-69, markers - 13 (differential dye), number of poliploid cells 11.0%

**DNA profile (STR):**

Amelogenin:	X,	X
CSF1PO:	9,	10
D13S317:	13.3,	13.3
D16S539:	9,	10
D5S818:	11,	12
D7S820:	8,	12
THO1:	7,	7
TPOX:	8,	12
vWA:	16,	18

**Plating efficiency:** 14% (ATCC)

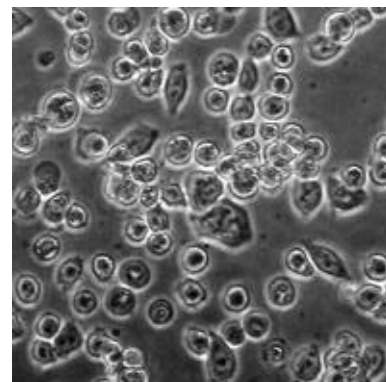
**Tumorigenicity:** non tumorigenic

**Other properties:** virus susceptibility: poliovirus type 1, adenovirus type 5, vesicular stomatitis (Indiana).

Isoenzymes G6PD, A

**Applications:** virology, toxicology, enzymology

**Collections:** ATCC CCL 2.2; ECACC 87110901; ICLC HTL 95020; SPBIC.



## HeLa TK<sup>-</sup>

**Origin:** human, epithelioid cervical carcinoma, strain of Hela. Submitted from Free University of Brussels, Belgium

Atlas of chromosomes of human and animal cell lines, S.E. Mamaeva, 2002. Moscow, Scientific world.

**Morphology:** epithelial-like

**Mode of cultivation:** monolayer

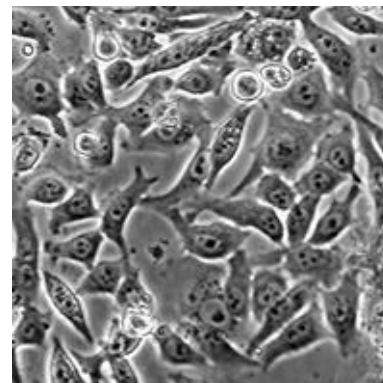
**Conditions for cultivation:** medium - DMEM

serum - FBS 10%

subculture procedure - cells detach from flask using trypsin 0.25%: EDTA 0.02% (1:3), split ratio 1:3 - 1:5, optimal

population density  $1.0-5.0 \times 10^4$  cells/cm<sup>2</sup>

cryoconservation - growth medium, 10% DMSO,  $1.5 \times 10^6$  cells/ml in ampule



**Viability after cryoconservation:** 90% (0 passage, dye trypan blue)

**Sterility:** tests for bacteria, fungi and mycoplasma were negative

**Species:** karyological and isoenzymological (LDH, G6PD) analysis

**Karyology:**  $2n=46$ , variability in the range between 57-61 chromosomes, modal number of chromosomes 60, markers - 22 (differential dye), number of poliploid cells 14.0%

**DNA profile (STR):**

Amelogenin:	X,	X
CSF1PO:	9,	10
D13S317:	13.3,	13.3
D16S539:	10,	10
D5S818:	11,	12
D7S820:	8,	12
THO1:	7,	7
TPOX:	8,	12
vWA:	16,	18

**Other properties:** deficient in thymidine kinase, resistant to 5-bromodeoxyuridine.

**Applications:** somatic cell genetics, cell biology

**Collections:** SPBIC.

**Origin:** human, hepatocyte carcinoma

Nature 1979. 282: 615-616; Science 1980. 209: 497-499.

**Morphology:** epithelial-like

**Mode of cultivation:** monolayer

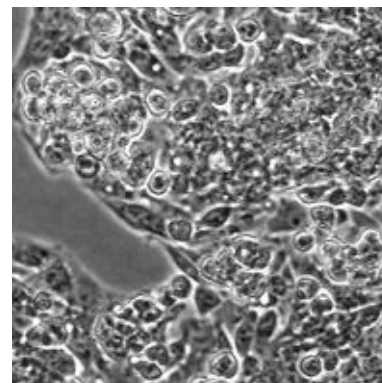
**Conditions for cultivation:** medium - EMEM, DMEM  
serum - FBS 10%

other components - NEAA 1%(EMEM),  
sodium pyruvate 0.1%

subculture procedure - cells detach from  
flask using trypsin 0.25%: EDTA 0.02%  
(1:3), split ratio 1:3 - 1:6, optimal

population density  $2.0-3.0 \times 10^4$  cells/cm<sup>2</sup>

cryoconservation - growth medium, 10%  
DMSO,  $1.0 \times 10^6$  cells/ml in ampule



**Viability after cryoconservation:** 98% (0 passage, dye trypan blue)

**Sterility:** tests for bacteria, fungi and mycoplasma were negative

**Species:** karyological and isoenzymological (LDH, G6PD) analysis

**Karyology:**  $2n=46$ , variability in the range between 49-57 chromosomes, modal  
number of chromosomes 55, number of polyploid cells - 5.6%.

**DNA profile (STR):**

Amelogenin:	X,	Y
CSF1PO:	10,	11
D13S317:	9,	13
D16S539:	12,	13
D5S818:	11,	12
D7S820:	10,	10
THO1:	9,	9
TPOX:	8,	9
vWA:	17,	17

**Tumorigenicity:** non tumorigenic in nude mice

**Other properties:** produce  $\alpha$ -fetoprotein, albumin,  $\alpha$ 2-macroglobulin,  $\alpha$ 1-antitrypsin,  
transferrin,  $\alpha$ 1-antichymotrypsin, haptoglobin, ceruloplasmin, plasminogen, complement  
(C3, C4), C3 activator, fibrinogen,  $\alpha$ 1-acid glycoprotein,  $\alpha$ 2-HS glycoprotein,  $\beta$ -  
lipoprotein, retinol binding protein.

**Applications:** biotechnology, biochemistry, virology, receptor study, enzymology,  
differentiation, cell biology

**Collections:** ATCC HB 8065; ECACC 85011430; SPBIC.

**Origin:** human, peripheral blood, promyelocytic leukemia.

Nature 1977. 270: 347-349; Blood 1979. 54: 713-733; Cytology (Russ.) 1992. 34: 123.  
Atlas of chromosomes of human and animal cell lines,  
S.E. Mamaeva, 2002. Moscow, Scientific world.

**Morphology:** lymphoblast-like

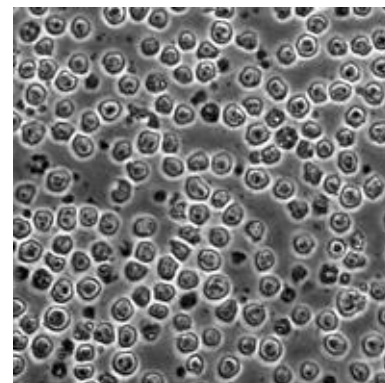
**Mode of cultivation:** suspension

**Conditions for cultivation:** medium - RPMI 1640 (Initial growth is sometimes by using Iscove's DMEM)

serum - FBS 20%

subculture procedure - split ratio 1:2,  
optimal population density  $1.0-5.0 \times 10^5$   
cells/cm<sup>2</sup>

cryoconservation - growth medium,  
5%DMSO,  $3.0-5.0 \times 10^6$  cells/ml in ampule



**Viability after cryoconservation:** 80% (0 passage, dye trypan blue)

**Sterility:** tests for bacteria, fungi and mycoplasma were negative

**Species:** karyological and isoenzymological (LDH, G6PD) analysis

**Karyology:**  $2n=46$ , variability in the range between 43-47 chromosomes, modal number of chromosomes 45, number of markers - 7 (differential dye), double minute chromosomes, number of polyploid cells 3%.

**DNA profile (STR):**

Amelogenin:	X,	X
CSF1PO:	13,	14
D13S317:	8,	11
D16S539:	11,	11
D5S818:	12,	12
D7S820:	11,	12
THO1:	7,	8
TPOX:	8,	11
vWA:	16,	16

**Plating efficiency:** the cells cannot be plated.

**Tumorigenicity:** tumorigenic in nude mice

**Other properties:** virus susceptibility: HIV-1, HTLV-1.

Isoenzymes G6PD, B; PGM1,1; PGM3,1; ES D,1; Me-2,1; AK 1,1; GLO-1,1.

Erythrocyte rosette tests: E, 4%; EA, 17%; EAC, 1%.

**Applications:** differentiation, pharmacodynamics, Tumorigenicity:

**Collections:** : ATCC CCL 240; ECACC 88112501; DSM ACC 3; ICLC HTL 95010; SPBIC.

**Origin:** human, kidney hypernephroma.

Biolog.Nauki 1985, 6: 29.

**Morphology:** epithelial-like

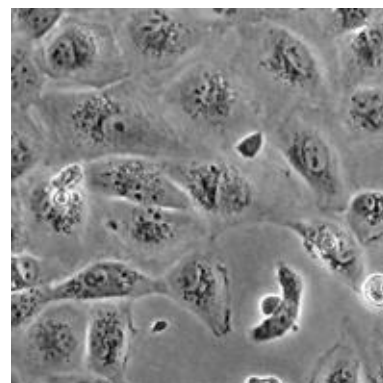
**Mode of cultivation:** monolayer

**Conditions for cultivation:** medium - DMEM

serum - FBS 10%

subculture procedure - cells detach from flask using trypsin 0.25%: EDTA 0.02% (1:3), split ratio 1:2 - 1:3.

cryoconservation - growth medium, 5-10% DMSO,  $1.0-1.5 \times 10^6$  cells/ml in ampule



**Viability after cryoconservation:** 75% (0 passage, dye trypan blue)

**Sterility:** tests for bacteria, fungi and mycoplasma were negative

**Species:** karyological analysis

**Karyology:**  $2n=46$ , variability in the range between 55-74 chromosomes, modal number of chromosomes 62.

**DNA profile (STR):**

Amelogenin:	X,	Y
CSF1PO:	11,	11
D13S317:	не установлено	
D16S539:	11,	12
D5S818:	12,	12
D7S820:	9,	11
THO1:	6,	9.3
TPOX:	8,	11
vWA:	15,	16, 17

**Tumorigenicity:** produce tumors in the cheek pouch of the hamster

**Other properties:** virus susceptibility: vesicular stomatitis, herpes simplex, cytomegalovirus, adenoviruses, RSV, encephalomyocarditis, parainfluenza 1 and 2, SV-40.

**Applications:** biochemistry, immunology, cell biology, virology.

**Collections:** SPBIC.



## Hos (TE85, clone F5)

**Origin:** human, osteosarcoma.

Cancer 1971. 27: 397-402; Int.J.Cancer 1975. 15: 23-29; Int.J.Cancer 1975. 16: 840-849. Atlas of chromosomes of human and animal cell lines, S.E. Mamaeva, 2002. Moscow, Scientific world.

**Morphology:** epithelial-like

**Mode of cultivation:** monolayer

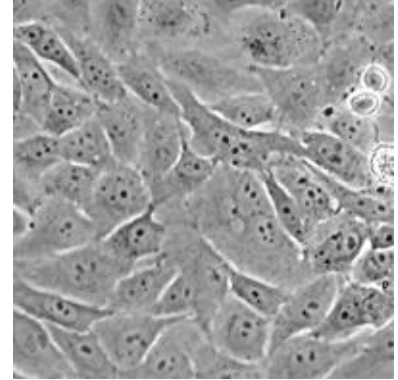
**Conditions for cultivation:** medium - DMEM

serum - FBS 10%

subculture procedure - cells detach from flask using trypsin 0.25%: EDTA 0.02%

(1:3), split ratio 1:2 - 1:6, optimal population density  $2.0-4.0 \times 10^4$  cells/cm<sup>2</sup>

cryoconservation - growth medium, 10% DMSO,  $2.5 \times 10^6$  cells/ml in ampule



**Viability after cryoconservation:** 95% (0 passage, dye trypan blue)

**Sterility:** tests for bacteria, fungi and mycoplasma were negative

**Species:** karyological and isoenzymological (LDH, G6PD) analysis

**Karyology:**  $2n=46$ , modal number of chromosomes 50, number of markers 12 (differential dye), number of polyploid cells 3.6%.

**DNA profile (STR):**

Amelogenin:	X,	X
CSF1PO:	12,	12
D13S317:	12,	12
D16S539:	10,	13
D5S818:	13,	13
D7S820:	11,	12
THO1:	6,	6
TPOX:	8,	11
vWA:	18,	18

**Other properties:** cells are sensitive to both virus and chemical transformation

**Applications:** virology, transformation, biochemistry

**Collections:** ATCC CRL 1543; ECACC 87070202; MWIHW; SPBIC.

**Origin:** human, ductal breast carcinoma

J.Natl.Cancer Inst. 1977. 58: 1795-1806.

**Morphology:** epithelial-like

**Mode of cultivation:** monolayer

**Conditions for cultivation:** medium - DMEM

serum - FBS 10%

other components - bovine insulin 10 $\mu$ g/ml

subculture procedure - cells detach from

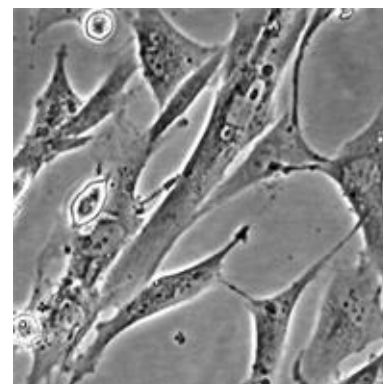
flask using trypsin 0.25%: EDTA 0.02%

(1:3), split ratio 1:3 - 1:5, optimal

population density 2.0-4.0 $\times 10^4$  cells/cm<sup>2</sup>

cryoconservation - growth medium, 10%

DMSO, 1.0 $\times 10^6$  cells/ml in ampule



**Viability after cryoconservation:** 90% (0 passage, dye trypan blue)

**Sterility:** tests for bacteria, fungi and mycoplasma were negative

**Species:** karyological analysis

**Karyology:** 2n= 46, variability in the range between 50-77 chromosomes, modal number of chromosomes 59, number of markers - 10 (differential dye), number of polyploid cells 15.8%.

**DNA profile (STR):**

Amelogenin:	X,	X
CSF1PO:	13,	13
D13S317:	11,	11
D16S539:	9,	12
D5S818:	11,	11
D7S820:	10,	10
THO1:	9,	9.3
TPOX:	8,	8
vWA:	17,	17

**Tumorigenicity:** tumorigenic in immunosuppressed mice

**Other properties:** estrogen receptors were not detected.

Isoenzymes G6PD, B; PGM<sub>1</sub>,1; PGM<sub>3</sub>,1; ES D,1; Me-2, 0; AK 1,1; GLO-1,1.

**Applications:** antitumor tests, radiotherapy, tumorigenicity:

**Collections:** ATCC HTB 126; ECACC 86082104; SPBIC.

**Origin:** human, fibrosarcoma.

Cancer 1974. 33: 1027-1033.

**Morphology:** epithelial-like

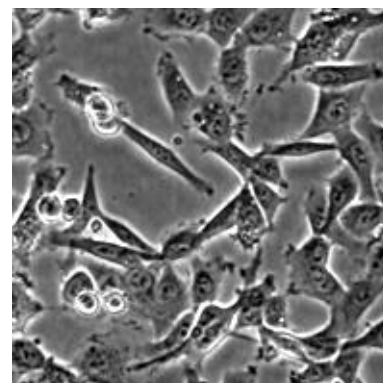
**Mode of cultivation:** monolayer

**Conditions for cultivation:** medium - DMEM

serum - FBS 10%

subculture procedure - cells detach from flask using trypsin 0.25%: EDTA 0.02% (1:3), split ratio 1:4 - 1:8, optimal population density  $1.0-4.0 \times 10^4$  cells/cm<sup>2</sup>

cryoconservation - growth medium, 5%DMSO,  $1.2 \times 10^6$  cells/ml in ampule



**Viability after cryoconservation:** 96% (0 passage, dye trypan blue)

**Sterility:** tests for bacteria, fungi and mycoplasma were negative

**Species:** isoenzymological (LDH, G6PD) analysis (LDH and G6PD)

**Karyology:**  $2n = 46$ , variability in the range between 44-48 chromosomes, modal number of chromosomes 46, pseudodiploid, about 40% of the cells had rearranged karyotypes.

**Plating efficiency:** 3%.

**DNA profile (STR):**

Amelogenin:	X,	Y
CSF1PO:	12,	12
D13S317:	12,	14
D16S539:	9,	12
D5S818:	11,	13
D7S820:	9,	10
THO1:	6,	6
TPOX:	8,	8
vWA:	14,	19

**Tumorigenicity:** tumorigenic in NIH Swiss mice immunosuppressed with anti-thymocytic serum.

**Other properties:** virus susceptibility: - RNA tumor viruses (RD 114, FeIV), poliovirus 1, vesic. stomatitis (Indiana).

Isoenzymes G6PD, B.

Chemotaxis, chemoinvasion, matrigel invasion.

Collagen production

**Applications:** molecular and cell biology, cytotoxicity, tumorigenicity, virology.

**Collections:** ATCC CCL 121; ECACC 85111505; SPBIC.

**Origin:** human, duodenum, adenocarcinoma

**Morphology:** epithelial-like

**Mode of cultivation:** monolayer

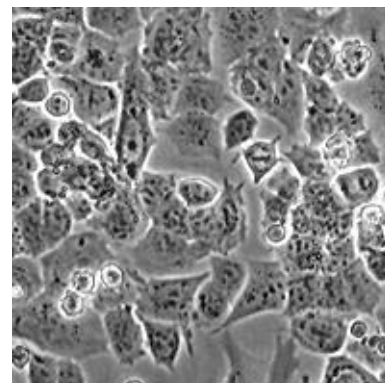
**Conditions for cultivation:** medium - EMEM

serum - FBS 10%

other components - NEAA 1%

subculture procedure - cells detach from flask using trypsin 0.25%: EDTA 0.02% (1:1 - 1:3), split ratio 1:3

cryoconservation - growth medium, 5-10% DMSO,  $1.0 \times 10^6$  cells/ml in ampule



**Viability after cryoconservation:** 90% (0 passage dye trypan blue)

**Sterility:** tests for bacteria, fungi and mycoplasma were negative

**Species:** isoenzymological (LDH, G6PD) analysis

**Karyology:**  $2n=46$ , variability in the range between 42-48 chromosomes, modal number of chromosomes 46, pseudodiploid, number of polyploid cells 0.4%, number of markers - 3 (differential dye).

**DNA profile (STR):**

Amelogenin:	X,	Y
CSF1PO:	11,	13
D13S317:	8,	11
D16S539:	10,	11
D5S818:	12,	13
D7S820:	9,	11
THO1:	7,	7
TPOX:	9,	11
vWA:	16,	18

**Tumorigenicity:** tumorigenic in nude mice

**Other properties:** isoenzymes PGM<sub>3,1-2</sub>; PGM<sub>1,1-2</sub>; ES D,1; Me-2,2; AK 1,1; GLO-1,2; G6PD,B

**Applications:** tumorigenicity, cell biology

**Collections:** ATCC HTB 40; SPBIC.

**Origin:** human, bone marrow, myeloma

Ann NY Acad.Sci. 1972. 190: 221-234; PNAS 1974. 71: 84-88; Nature 1974. 251: 443-444; J.Biol. Chem. 1974. 249: 1661-1667; J.Biol. Chem. 1976. 251: 6844-6851; Atlas of chromosomes of human and animal cell lines, S.E. Mamaeva, 2002. Moscow, Scientific world.

**Morphology:** lymphoblast-like

**Mode of cultivation:** suspension

**Conditions for cultivation:** medium - RPMI 1640

serum - FBS 10%

subculture procedure - optimal

population density  $2.0-4.0 \times 10^5$  cells/ml

cryoconservation - growth medium, 10%

DMSO,  $3.0-4.0 \times 10^6$  cells/ml in ampule

**Viability after cryoconservation:** 90% (0 passage, dye trypan blue)

**Sterility:** tests for bacteria, fungi and mycoplasma were negative

**Species:** karyological and isoenzymological (LDH, G6PD) analysis

**Karyology:**  $2n=46$ , variability in the range between 43-48 chromosomes, modal number of chromosomes 46, normal human karyotype (46, XX), but heterochromatin areas of the two homologue chromosomes 1 – decondensation, number of polyploid cells 7.5%.

**DNA profile (STR):**

Amelogenin:	X,	X
CSF1PO:	10,	11
D13S317:	9,	11
D16S539:	9,	13
D5S818:	13,	13
D7S820:	11,	12
THO1:	6,	9.3
TPOX:	11,	11
vWA:	14,	17

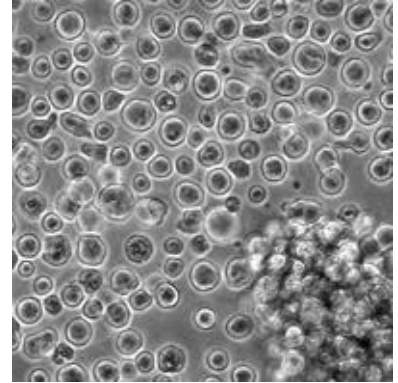
**Plating efficiency:** the cells cannot be plated

**Other properties:** isoenzymes PGM<sub>1,1-2</sub>; PGM<sub>3</sub>, 0; ES T-D,1; Me-2, 2; GLO-1,1-2; G6PD, B.

Human growth hormone receptor, insulin receptor, calcitonin receptor. Erythrocyte rosette tests: E, 1%; EA, 0; EAC, 13%.

**Applications:** biotechnology (Ig G kappa production), endocrinology, Tumorigenicity:

**Collections:** ATCC CCL 159; DSM ACC 117; ECACC 86051302; SPBIC.



**Origin:** human, neuroblastoma

Cancer Res. 1970. 30: 2110. Atlas of chromosomes of human and animal cell lines, S.E. Mamaeva, 2002. Moscow, Scientific world.

**Morphology:** fibroblast- and neuroblast-like

**Mode of cultivation:** monolayer

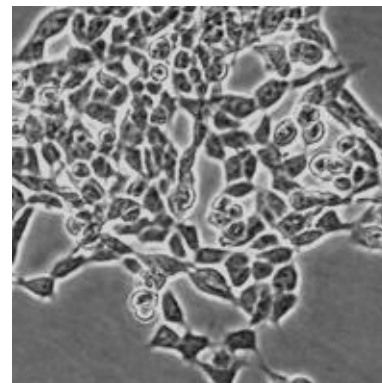
**Conditions for cultivation:** medium - DMEM

serum - FBS 10%

subculture procedure - cells detach from flask using trypsin 0.25%: EDTA 0.02% (1:3), split ratio 1:3 - 1:6, optimal

population density  $2.0-3.0 \times 10^4$  cells/cm<sup>2</sup>

cryoconservation - growth medium, 10% DMSO,  $1.5-2.0 \times 10^6$  cells/ml in ampule



**Viability after cryoconservation:** 76% (0 passage, dye trypan blue)

**Sterility:** tests for bacteria, fungi and mycoplasma were negative

**Species:** karyological and isoenzymological (LDH, G6PD) analysis

**Karyology:**  $2n=46$ , variability in the range between 42-51 chromosomes, modal number of chromosomes 48, number of markers - 2 (differential dye), number of polyploid cells 16%.

**DNA profile (STR):**

Amelogenin:	X,	Y
CSF1PO:	11,	12
D13S317:	9,	9
D16S539:	8,	8
D5S818:	11,	12
D7S820:	9,	10
THO1:	7,	9.3
TPOX:	11,	11
vWA:	15,	15

**Plating efficiency:** less than 1%.

**Tumorigenicity:** tumorigenic in nude mice

**Other properties:** virus susceptibility: vesicular stomatitis (Indiana), herpes simplex, vaccinia, adenovirus 12, Coxsackie B3.

Isoenzymes G6PD, B;  
neurotransmitter synthesis.

**Applications:** tumorigenicity, immunology, differentiation, electrophysiology, cell biology

**Collections:** ATCC CCL 127; ECACC 86041809; ICLC HTL 96021; SPBIC.

**Origin:** human, T-lymphoblastic leukemia

Submitted from Institute of Immunology, Moscow. Atlas of chromosomes of human and animal cell lines, S.E. Mamaeva, 2002. Moscow, Scientific world.

**Morphology:** lymphoblast-like

**Mode of cultivation:** suspension

**Conditions for cultivation:** medium - RPMI 1640

serum - FBS 10%

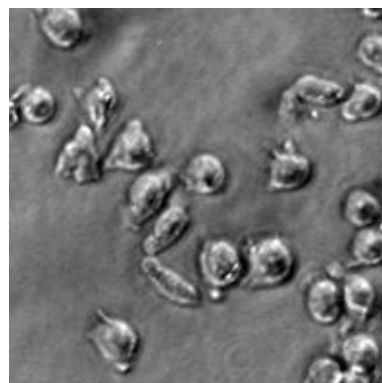
subculture procedure - optimal

population density  $3.0-9.0 \times 10^5$  cells/ml

cryoconservation - growth medium,

10% DMSO,  $3.0-4.0 \times 10^6$  cells/ml in

ampule



**Viability after cryoconservation:** 90% (0 passage, dye trypan blue)

**Sterility:** tests for bacteria, fungi and mycoplasma were negative

**Species:** karyological and isoenzymological (LDH, G6PD) analysis

**Karyology:**  $2n=46$ , variability in the range between 41-49 chromosomes, modal number of chromosomes 46-47, number of markers - 2 (differential dye).

**DNA profile (STR):**

Amelogenin:	X,	Y
CSF1PO:	11,	12
D13S317:	8,	11
D16S539:	11,	11
D5S818:	9,	9
D7S820:	8,	10
THO1:	6,	9.3
TPOX:	8,	10
vWA:	18,	18

**Other properties:** IL-2 synthesis, T-cell marker CD 3.

**Applications:** immunology, biochemistry, differentiation

**Collections:** SPBIC.

**Origin:** human, chronic myelogenous leukemia (pleural effusion).

Blood 1975. 45: 321-334; J.Natl.Cancer Inst. 1977. 59; 77; Int.J.Cancer 1979. 23: 143-147; Leukemia Res. 1979. 3; 363; Proc. 37<sup>th</sup> Ann.Meet.Electron Microsc.Soc.Amer, tex. 1979: 234; Blood 1980. 56: 344-350; J.Biol.Chem. 1980. 255: 3266; Biochem.J. 1981. 193: 361; Proc.Soc.Exp.Biol.Med. 1981. 166: 546-550; J.Immunol. 1982. 129; 2504; Exp.Hematol. 1983. 11: 601-610; Clin.Haematol.1984.13:461; Biology of the cell in culture. L. Nauka,1984.279. Atlas of chromosomes of human and animal cell lines, S.E. Mamaeva, 2002. Moscow, Scientific world.

**Morphology:** erythromyeloblastoid

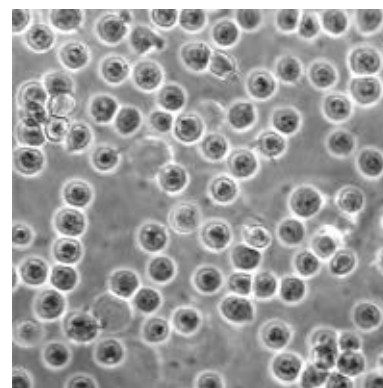
**Mode of cultivation:** suspension

**Conditions for cultivation:** medium - RPMI 1640

serum - FBS 10%

subculture procedure - optimal  
population density  $1.0 \times 10^5$ - $1.0 \times 10^6$   
cells/ml

cryoconservation - growth medium,  
10% DMSO,  $3.0$ - $7.0 \times 10^6$  cells/ml in  
ampule



**Viability after cryoconservation:** 93% (0 passage, dye trypan blue)

**Sterility:** tests for bacteria, fungi and mycoplasma were negative

**Species:** karyological and isoenzymological (LDH, G6PD) analysis

**Karyology:**  $2n=46$ , there are some sublines K-562 with different karyotypic structure.

One from sublines is: variability in the range between 55-69 chromosomes, modal number of chromosomes 66, number of markers - 12 (differential dye), number of polyploid cells 3%.

<b>DNA profile (STR):</b>	Amelogenin:	X,	X
	CSF1PO:	9,	10
	D13S317:	8,	8
	D16S539:	11,	12
	D5S818:	11,	12
	D7S820:	9,	11
	THO1:	9.3,	9.3
	TPOX:	8,	9
	vWA:	16,	16

**Plating efficiency:** the cells cannot be plated

**Tumorigenicity:** tumorigenic in nude mice

**Other properties:** haemoglobin synthesis.

Isoenzymes AK 1,1; ES D,1; GLO-1, 2; G6PD, B; PGM<sub>1</sub>, 0; PGM<sub>3</sub>,1; Me-2,0.

Erythrocyte rosette tests: E, 1%; EA, 34%; EAC, 2%.

Capable to differentiate into progenitors of the erythrocytic, granulocytic and monocytic series.

Not contained B- and T-markers.

**Applications:** differentiation, cell biology, natural killer assay, pharmacodynamics.

**Collections:** ATCC CCL 243; ECACC 89121407; DSM ACB 10; ICLC HTL 94001; MWIIW; SPBII; SPBIC.



**Origin:** human, acute myelogenous leukemia (bone marrow)  
 Science 1978. 200: 1153-1154; Blood 1980. 56: 344-350; Blood 1979. 54: Suppl. 1, 174a.

**Morphology:** myeloblastoid

**Mode of cultivation:** suspension

**Conditions for cultivation:** medium - RPMI 1640

serum - FBS 20%

subculture procedure - optimal

population density  $3.0-9.0 \times 10^5$  cells/ml

cryoconservation - growth medium, 5%

DMSO,  $3.0-4.0 \times 10^6$  cells/ml in ampule

**Viability after cryoconservation:** 85% (0 passage, dye trypan blue)

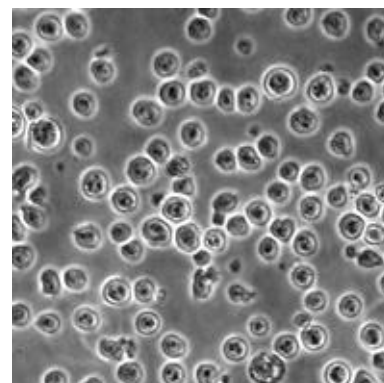
**Sterility:** tests for bacteria, fungi and mycoplasma were negative

**Species:** karyological and isoenzymological (LDH, G6PD) analysis

**Karyology:**  $2n=46$ , variability in the range between 44-49 chromosomes, modal number of chromosomes 46-47, number of markers - 5 (differential dye) (ATCC).

**DNA profile (STR):**

Amelogenin:	X,	Y
CSF1PO:	7,	7
D13S317:	11,	12
D16S539:	10,	11
D5S818:	13,	13
D7S820:	8,	10
THO1:	7,	8
TPOX:	7,	9
vWA:	14,	19



**Plating efficiency:** the cells cannot be plated.

**Tumorigenicity:** non tumorigenic

**Other properties:** isoenzymes G6PD, B; PGM<sub>1</sub>, 1; PGM<sub>3</sub>, 0; ES D, 1; Me-2, 1; AK 1,0; GLO-1,2.

Have no surface Ig antigens.

Erythrocyte rosette tests: E, 0; EA, 2%; EAC, 0.

HLA cell line phenotype A 30, 31; B 35; Cw 4.

Express the human DR antigen.

Differentiation into non-dividing macrophages when exposed to phorbol esters; formation of colonies in soft-agar culture when exposed to colony-stimulating factor

**Applications:** tumorigenicity, differentiation

**Collections:** ATCC CCL 246; DSM ACC 14; ECACC 86111306; SPBIC.

## M-FetMSC

**Origin:** human, mesenchymal stem cells from Muscle of a limb of 5-6 week embryo.  
Tsitologiya. 2014. 56 (8): 562 – 573.

**Morphology:** fibroblast-like.

**Mode of cultivation:** monolayer

**Conditions for cultivation:** medium - DMEM/F12

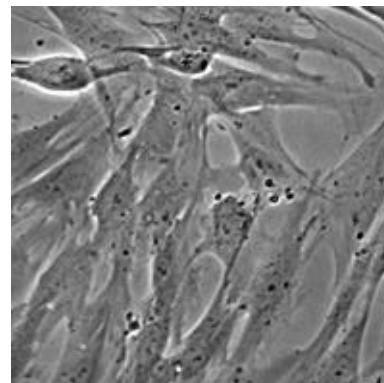
serum - FBS 10%

subculture procedure - cells detach from flask using trypsin 0.25%: EDTA 0.02% (1:3), split ratio: 1:3-1:5, optimal

population density  $4.0-5.0 \times 10^4$  cells/cm<sup>2</sup>

cryoconservation - growth medium, 10%

DMSO,  $1.5-2.0 \times 10^6$  cells/ml in ampule



**Viability after cryoconservation:** 90% (0 passage, dye trypan blue)

**Sterility:** tests for bacteria, fungi and mycoplasma were negative.

**Species:** karyological analysis.

**Karyology:**  $2n=46$ , modal number of chromosomes 46 ( $99.1 \pm 0.9\%$ ), normal human karyotype (46, XY), number of polyploid cells 2.2%.

**DNA profile (STR):**

Amelogenin:	X,	Y
CSF1PO:	9,	12
D13S317:	11,	12
D16S539:	11,	11
D5S818:	12,	13
D7S820:	10,	12
THO1:	7,	8
TPOX:	8,	11
vWA:	14,	15

**Other properties:** finite lifetime culture; average population doubling time 25.0 h; The presence of surface antigens specific for human MSCs: CD44, CD73, CD90, CD105, HLA-ABC, vimentin and the lack of CD34, HLA-DR; The ability to induced differentiation into osteogenic, chondrogenic and adipogenic directions. Induced skeletal-muscle differentiation with the formation of myotube and Zdisks.

**Applications:** cell biology, myogenesis, biotechnology, feeder for cultivation embryonic stem cells.

**Collections:** SPBIC.

**Origin:** human, breast adenocarcinoma (pleural effusion)  
J.Natl.Cancer Inst. 1973. 51: 1409-1416.

**Morphology:** epithelial-like

**Mode of cultivation:** monolayer

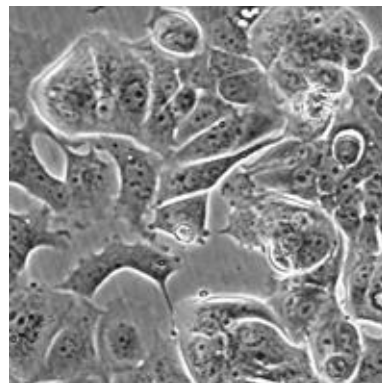
**Conditions for cultivation:** medium - EMEM

serum - FBS 10%

other components - NEAA 1%, bovine insulin 10  $\mu$ /ml.

subculture procedure - cells detach from flask using trypsin 0.25%: EDTA 0.02% (1:3), split ratio 1:3, optimal population density  $2.0-4.0 \times 10^4$  cells/cm<sup>2</sup>

cryoconservation - growth medium, 8-9%DMSO,  $1.0 \times 10^6$  cells/ml in ampule



**Viability after cryoconservation:** 94% (0 passage, dye trypan blue)

**Sterility:** tests for bacteria, fungi and mycoplasma were negative

**Species:** karyological analysis

**Karyology:**  $2n=46$ , variability in the range between 67-87 chromosomes, modal number of chromosomes 79-82, number of markers 2, large acrocentric and submetacentric chromosomes (routine dye), 29-34 (differential dye), number of polyploid cells 0.6%.

**DNA profile (STR):**

Amelogenin:	X,	X
CSF1PO:	10,	10
D13S317:	11,	11
D16S539:	11,	12
D5S818:	11,	12
D7S820:	8,	9
THO1:	6,	6
TPOX:	9,	12
vWA:	14,	15

**Tumorigenicity:** tumorigenic in nude mice

**Other properties:** isoenzymes PGM<sub>3</sub>, 1-2; PGM<sub>1</sub>, 2; ES D, 1; AK 1, 1; GLO-1, 1-2; G6PD, B. Estrogen receptor positive.

Estradiol synthesis.

Cells may carry B- or C-type virus.

The capability of forming domes.

**Applications:** receptor study, chemotherapeutic agents, tumorigenicity, cell biology, virology.

**Collections:** ATCC HTB 22; ECACC 86012803; ICLC HTL 95021; SPBIC.

**Origin:** human, osteosarcoma

Antimicrob. Agents Chemother. 1977. 12: 11-15., Atlas of chromosomes of human and animal cell lines, S.E. Mamaeva, 2002. Moscow, Scientific world.

**Morphology:** fibroblast-like

**Mode of cultivation:** monolayer

**Conditions for cultivation:** medium - EMEM

serum - FBS 10%

other components - NEAA 1%

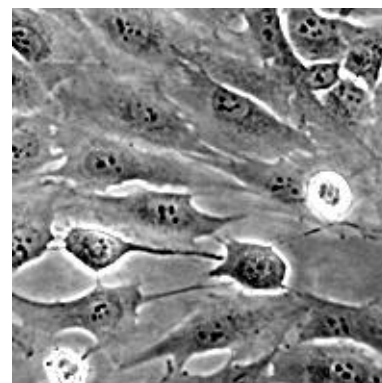
subculture procedure - cells detach from flask using trypsin 0.25%: EDTA 0.02%

(1:3), split ratio 1:3 - 1:6, optimal

population density  $2.0-4.0 \times 10^4$  cells/cm<sup>2</sup>

cryoconservation - growth medium, 10%

DMSO,  $1.0 \times 10^6$  cells/ml in ampule



**Viability after cryoconservation:** 90% (0 passage, dye trypan blue)

**Sterility:** tests for bacteria, fungi and mycoplasma were negative

**Species:** karyological analysis

**Karyology:**  $2n=46$ , variability in the range between 59-65 chromosomes, modal number of chromosomes 63, number of markers - 22 (differential dye), number of polyploid cells - 2%.

<b>DNA profile (STR):</b>	Amelogenin:	X,	Y
	CSF1PO:	10,	12
	D13S317:	11,	11
	D16S539:	11,	11
	D5S818:	11,	12
	D7S820:	10,	10
	THO1:	9.3,	9.3
	TPOX:	8,	11
	vWA:	16,	19

**Applications:** biotechnology (interferon production), cell biology

**Collections:** ATCC CRL 1427, ECACC 86051601; SPBIC.

## M-HeLa clone 11

**Origin:** human, epithelioid cervical carcinoma, strain of HeLa, clone of M-HeLa  
J.Exp.Med. 1953, 97: 695; Cytology (Russ) 1986, 28: 56 - 61

**Morphology:** epithelial-like

**Mode of cultivation:** monolayer

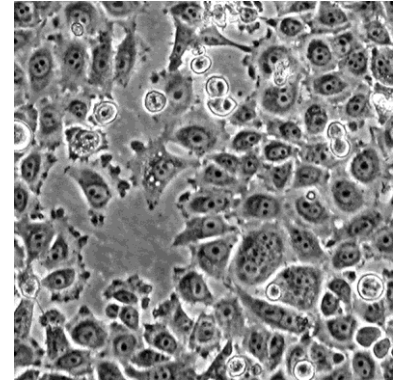
**Conditions for cultivation:** medium - EMEM

serum - FBS 10%

other components - NEAA 1%

subculture procedure - cells detachment using trypsin 0.25%: EDTA 0.02% (1:3), split ratio 1:3 - 1:6, optimal population density  $2.0-4.0 \times 10^4$  cells/cm<sup>2</sup>

cryoconservation - growth medium, 10% DMSO,  $1.0-2.0 \times 10^6$  cells/ml in ampule



**Viability after cryoconservation:** 80% (0 passage, dye trypan blue)

**Sterility:** bacteria, fungi and mycoplasma were negative

**Species specificity:** karyological and isoenzymological (LDH, G6PD) analysis

**Karyology:**  $2n=46$ , variability in the range between 49-50 chromosomes, modal number of chromosomes 50, number of markers - 13 (differential dye), number of polyploid cells - 2.4%.

**DNA profile (STR):**

Amelogenin:	X,	X
CSF1PO:	9,	10
D13S317:	13.3,	13.3
D16S539:	9,	10
D5S818:	11,	12
D7S820:	12,	12
THO1:	7,	7
TPOX:	8,	8
vWA:	16,	18

**Plating efficiency:** 60%

**Applications:** cell biology, tumorigenicity, virology

**Collections:** SPBIC.

**Origin:** human, pancreatic carcinoma  
Int.J.Cancer 1977. 19: 128-135.

**Morphology:** epithelial-like

**Mode of cultivation:** monolayer

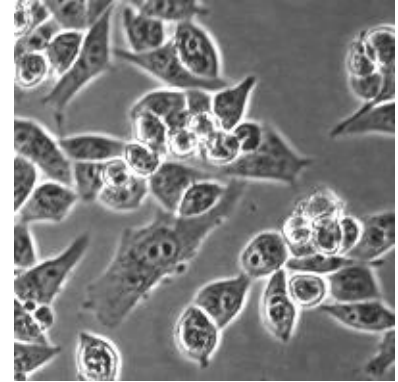
**Conditions for cultivation:** medium - DMEM

serum - FBS 10%+HS 2.5%

subculture procedure - cells detach from flask using trypsin 0.25%: EDTA 0.02% (1:3), split ratio 1:2 - 1:3, optimal

population density  $2.0-3.0 \times 10^4$  cells/cm<sup>2</sup>

cryoconservation - growth medium, 10% DMSO,  $3.0 \times 10^6$  cells/ml in ampule



**Viability after cryoconservation:** 90% (0 passage, dye trypan blue)

**Sterility:** tests for bacteria, fungi and mycoplasma were negative

**Species:** karyological analysis

**Karyology:**  $2n=46$ , modal number of chromosomes 61, number of markers - 16-20 (differential dye).

**DNA profile (STR):**

Amelogenin:	X,	X
CSF1PO:	10,	10
D13S317:	12,	13
D16S539:	10,	13
D5S818:	12,	13
D7S820:	12,	13
THO1:	9,	10
TPOX:	9,	9
vWA:	15,	15

**Other properties:** isoenzymes G6PD, B.

Sensitive to asparaginase

**Applications:** tumorigenicity, enzymology, cell biology

**Collections:** ATCC 3RL 1420; ECACC 85062806; SPBIC.

## MNNG-HOS (TE 85, clon F-5)

**Origin:** human, osteosarcoma, chemically transformed (MNNG 0.1  $\mu$ /ml)  
Nature 1975. 256: 51; Int.J.Cancer 1977. 19: 505.

**Morphology:** epithelial-like

**Mode of cultivation:** monolayer

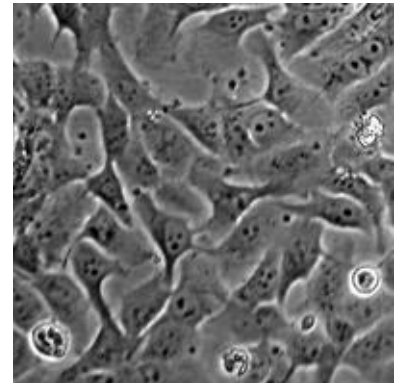
**Conditions for cultivation:** medium - DMEM

serum - FBS 10%

subculture procedure - cells detach from flask using trypsin 0.25%: EDTA 0.02% (1:3), split ratio 1:4 - 1:6, optimal

population density  $2.0-4.0 \times 10^4$  cells/cm<sup>2</sup>

cryoconservation - growth medium, 10% DMSO,  $1.0-2.0 \times 10^6$  cells/ml in ampule



**Viability after cryoconservation:** 98% (0 passage, dye trypan blue)

**Sterility:** tests for bacteria, fungi and mycoplasma were negative

**Species:** karyological and isoenzymological (LDH, G6PD) analysis

**Karyology:**  $2n=46$ , variability in the range between 63-74 chromosomes, modal number of chromosomes 69-70, number of polyploid cells 2.2%.

**DNA profile (STR):**

Amelogenin:	X,	X
CSF1PO:	12,	12
D13S317:	12,	12
D16S539:	10,	13
D5S818:	13,	13
D7S820:	11,	12
THO1:	6,	6
TPOX:	8,	11
vWA:	18,	18

**Tumorigenicity:** tumorigenic in nude mice

**Applications:** tumorigenicity, transformation

**Collections:** ATCC CRL 1547; ECACC 87070201; SPBIC.

## MOLT-3

**Origin:** human, T-lymphoblastic leukemia, peripheral blood.

J.Natl.Cancer Inst. 1972. 49: 891-895., Atlas of chromosomes of human and animal cell lines, S.E. Mamaeva, 2002. Moscow, Scientific world.

**Morphology:** lymphoid

**Mode of cultivation:** suspension

**Conditions for cultivation:** medium - RPMI 1640

serum - FBS 10%

subculture procedure - optimal

population density  $5.0-6.0 \times 10^5$  cells/ml

cryoconservation - growth medium,

10% DMSO,  $5.0 \times 10^6$  cells/ml in ampule

**Viability after cryoconservation:** 80% (0 passage, dye trypan blue)

**Sterility:** tests for bacteria, fungi and mycoplasma were negative

**Species:** karyological and isoenzymological (LDH, G6PD) analysis

**Karyology:**  $2n=46$  - modal number of chromosomes 98, number of markers - 4 (differential dye), number of polyploid cells 1.0%.

**DNA profile (STR):**

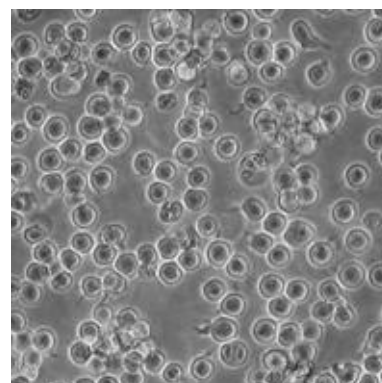
Amelogenin:	X,	Y
CSF1PO:	11,	12
D13S317:	12,	13
D16S539:	11,	14
D5S818:	12,	12
D7S820:	8,	10
THO1:	6,	8
TPOX:	8,	8
vWA:	17,	17

**Other properties:** virus susceptibility: HIV.

The cells form rosettes with sheep erythrocytes.

**Applications:** tumorigenicity, virology

**Collections:** ATCC CRL 1552; DSM ACC 84; ECACC 90021901; SPBIC.





**Origin:** human, T-lymphoblastic leukemia, peripheral blood.

J.Natl.Cancer Inst. 1972. 49: 891-895; J.Immunol. 1982. 129: 2504-2510;  
Int.J.Immunopharmacol. 1988. 10: 907-911; Glukhova L.A. PhD Thesis; SPBIC,  
St.Petersburg. 1992.

**Morphology:** lymphoblast-like

**Mode of cultivation:** suspension

**Conditions for cultivation:** medium - RPMI 1640

serum - FBS 10%

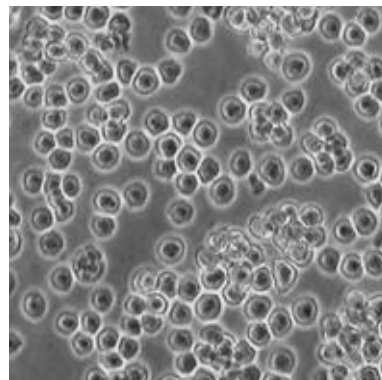
subculture procedure - optimal

population density  $2.0-5.0 \times 10^5$  cells/ml

cryoconservation - growth medium,

10% DMSO,  $5.0 \times 10^6$  cells/ml in

ampule



**Viability after cryoconservation:** 94% (0 passage, dye trypan blue)

**Sterility:** tests for bacteria, fungi and mycoplasma were negative

**Species:** karyological and isoenzymological (LDH, G6PD) analysis

**Karyology:**  $2n=46$ , variability in the range between 77-100 chromosomes, modal number of chromosomes 97, number of markers - 6 (differential dye), number of polyploid cells 2.0%.

**DNA profile (STR):**

Amelogenin:	X,	Y
CSF1PO:	11,	12, 13
D13S317:	12,	13
D16S539:	11,	14
D5S818:	11,	12
D7S820:	8,	10, 11
THO1:	6,	8
TPOX:	8,	8
vWA:	17,	18

**Tumorigenicity:** tumorigenic in nude mice

**Other properties:** virus susceptibility: measles,  $\alpha$ -viruses

Terminal deoxynucleotidyl transferase activity is high.

The cells form rosettes with sheep erythrocytes.

**Applications:** biochemistry, cytotoxicity, differentiation, virology, tumorigenicity, immunology

**Collections:** ATCC CRL 1582; ECACC 85011413; MWIIV; SPBIC.

**Origin:** human, mesenchymal stem cells from pulp of a deciduous tooth of a child. Tsitologiya. 2018. 60 (12): 955 – 268.

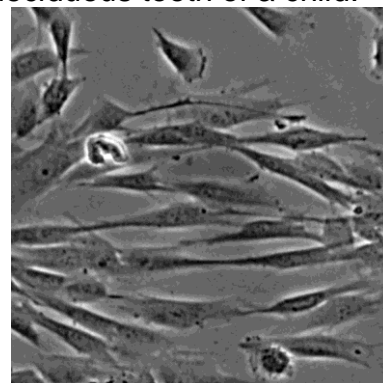
**Morphology:** fibroblast-like

**Mode of cultivation:** monolayer

**Conditions for cultivation:** medium – DMEM/F12  
serum - FBS 10%

subculture procedure - cells detach from flask using trypsin 0.25%: EDTA 0.02% (1:3), split ratio: 1:3-1:4, optimal population density 2.0- 4.0x10<sup>4</sup> cells/cm<sup>2</sup>

cryoconservation - growth medium, 10% DMSO, 1.0-1.5x10<sup>6</sup> cells/ml in ampule.



**Viability after cryoconservation:** 90% (0 passage, dye trypan blue)

**Sterility:** tests for bacteria, fungi and mycoplasma were negative.

**Species:** karyological analysis.

**Karyology:** 2n=46, modal number of chromosomes 46 (99.0 ± 1.0 %), normal human karyotype (46, XX), number of poliploid cells 7.8 %.

**DNA profile (STR):**

Amelogenin:	X,	X
CSF1PO:	11,	11
D13S317:	8,	9
D16S539:	11,	11
D5S818:	9,	11
D7S820:	8,	10 12
THO1:	6,	8 9.3
TPOX:	8,	11
vWA:	15,	16 17

**Plating efficiency:** 32.8%.

**Other properties:** finite lifetime culture; average population doubling time 32.8 h.

The presence of surface antigens specific for human MSCs: CD44, CD73, CD90, CD105, HLA-ABC and the lack of CD34, HLA-DR;

The ability to induced differentiation into osteogenic and chondrogenic directions; the expression of neuronal differentiation gene.

**Applications:** cell biology, biotechnology.

**Collections:** SPBIC.

**Origin:** human, pulpa of tooth of six year old boy (mesenchymal stem cells)

Tsitologiya. 2023. 65 (5): 420-436.

**Morphology:** fibroblast-like

**Mode of cultivation:** monolayer

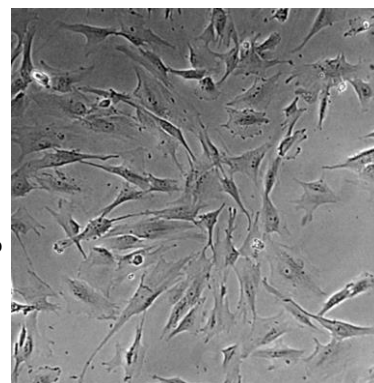
**Conditions for cultivation:**

medium – DMEM/F12

serum - FBS 10%

subculture procedure - cells detach from flask using 0.25% solution trypsin:EDTA, split ratio: 1:3-1:4, optimal population density  $4.0-5.0 \times 10^3$  cells/cm<sup>2</sup>

cryoconservation - growth medium, 10% DMSO,  $2.5 \times 10^6$  cells/ml in ampule.



**Viability after cryoconservation:** 83% (0 passage, dye trypan blue)

**Sterility:** tests for bacteria, fungi and mycoplasma were negative.

**Species:** karyological analysis.

**Karyology:**  $2n=46$ , modal number of chromosomes 46 ( $98.0 \pm 1.4$  %), normal human karyotype (XY), number of poliploid cells  $4.6 \pm 0.7$ %.

**DNA profile (STR):**

Amelogenin:	X,	Y
CSF1PO:	11,	13
D13S317:	9,	13
D16S539:	11,	12
D5S818:	9,	11
D7S820:	8,	10
THO1:	6,	7
TPOX:	8,	11
vWA:	15,	16

**Plating efficiency:**  $15.3 \pm 1.8$ %.

**Other properties:** finite lifetime culture; at the 6 passage the average time of one doubling of the cell population is  $26.6 \pm 0.32$  h. The time of the active (logarithmic) growth phase is 96 h. The phase of active replicative senescence occurs at 32 passage. The presence of surface antigens specific for human MSCs: CD44, CD73, CD90, CD105, vimentin, HLA-ABC and the lack of CD34, CD45, HLA-DR; The ability to induced differentiation into osteogenic, adipogenic and chondrogenic directions in early 6 passage but in the late 40 passage there is a decrease in the differentiation potential in the adipogenic directions.

**Applications:** cell biology, biotechnology.

**Collections:** SPBIC.

**Origin:** human, pulpa of tooth of six year old girl (mesenchymal stem cells)

Tsitologiya. 2023. 65 (5): 420-436.

**Morphology:** fibroblast-like

**Mode of cultivation:** monolayer

**Conditions for cultivation:**

medium – DMEM/F12

serum - FBS 10%

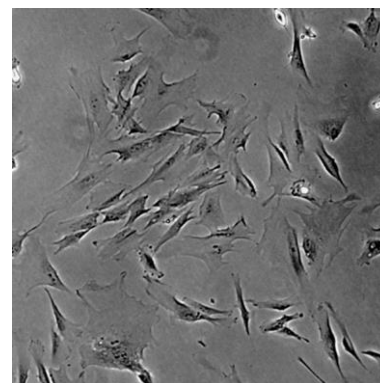
subculture procedure - cells detach from flask using 0.025

olution trypsin:EDTA, split ratio: 1:2 - 1:3, optimal

population density  $4.0- 5.0 \times 10^3$  cells/cm<sup>2</sup>

cryoconservation - growth medium, 10% DMSO,  $1.0 \times 10^6$

cells/ml in ampule.



**Viability after cryoconservation:** 87% (0 passage, dye trypan blue)

**Sterility:** tests for bacteria, fungi and mycoplasma were negative.

**Species:** karyological analysis.

**Karyology:**  $2n=46$ , modal number of chromosomes 46 ( $98.0 \pm 1.4$  %), normal human karyotype (XX), number of poliploid cells  $6.0 \pm 0.8$ %.

**DNA profile (STR):**

Amelogenin:	X,	X
CSF1PO:	10,	11
D13S317:	10,	10
D16S539:	11,	12
D5S818:	12,	13
D7S820:	11,	12
THO1:	9.3,	9.3
TPOX:	8,	11
vWA:	17,	19

**Plating efficiency:**  $16.6 \pm 2.8$ %.

**Other properties:** finite lifetime culture; at the 6 passage the average time of one doubling of the cell population is  $37.5 \pm 2.2$  h. The time of the active (logarithmic) growth phase is 72 h. The phase of active replicative senescence occurs at 16 passage.

The presence of surface antigens specific for human MSCs: CD44, CD73, CD90, CD105, HLA-ABC and the lack of CD34, CD45, HLA-DR;

The ability to induced differentiation into osteogenic and chondrogenic directions in early and later passages; reduced character adipogenic differentiation in the early passage; absence adipogenic differentiation in the later 18 passage.

**Applications:** cell biology, biotechnology.

**Collections:** SPBIC.

**Origin:** human, placenta (mesenchymal stem cells)

Tsitologiya. 2020. 62 (9): 713 – 727.

**Morphology:** fibroblast-like

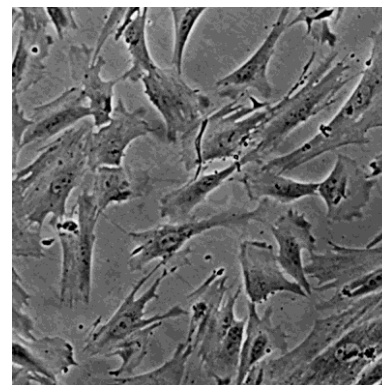
**Mode of cultivation:** monolayer

**Conditions for cultivation:** medium – DMEM/F12  
serum - FBS 10%

subculture procedure - cells detach from flask using trypsin 0.25%: EDTA 0.02% (1:3), split ratio: 1:3, optimal

population density  $4.0- 5.0 \times 10^3$  cells/cm<sup>2</sup>

cryoconservation - growth medium, 10% DMSO,  $2.5 \times 10^6$  cells/ml in ampule.



**Viability after cryoconservation:** 80% (0 passage, dye trypan blue)

**Sterility:** tests for bacteria, fungi and mycoplasma were negative.

**Species:** karyological analysis.

**Karyology:**  $2n=46$ , modal number of chromosomes 46 ( $98.0 \pm 1.4$  %), normal human karyotype (46, XX), number of polyploid cells  $4.0 \pm 0.6$ %.

**DNA profile (STR):**

Amelogenin:	X,	X
CSF1PO:	11,	14
D13S317:	8,	11
D16S539:	11,	13
D5S818:	11,	12
D7S820:	10,	12
THO1:	6,	9.3
TPOX:	11,	11
vWA:	17	18

**Plating efficiency:** 0.0 %.

**Other properties:** finite lifetime culture; at the 6 passage the average time of one doubling of the cell population is 14.5 h. The time of the active (logarithmic) growth phase is 24 h. The phase of active replicative senescence occurs at 16-18 passages. The logarithmic growth phase is 24 h. The average time of one doubling of the cell population is 44.1 h.

The presence of surface antigens specific for human MSCs: CD44, CD73, CD90, CD105, vimentin, HLA-ABC and the lack of CD34, CD45, HLA-DR;

The ability to induced differentiation into osteogenic, adipogenic and chondrogenic directions; but in the later passages there is a decrease in the differentiation potential in the osteogenic and adipogenic directions.

**Applications:** cell biology, biotechnology.

**Collections:** SPBIC.

**Origin:** human, mesenchymal stem cells from Wharton jelly of the umbilical cord. Tsitologiya. 2017. 59 (5): 315-327; Tsitologiya. 2017. 59 (9): 574-587.

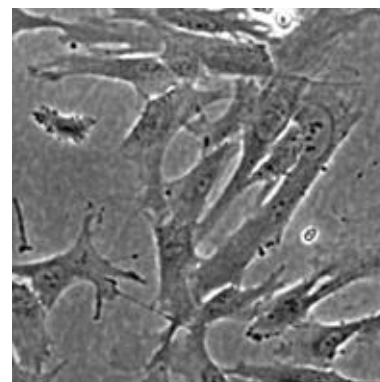
**Morphology:** fibroblast-like.

**Mode of cultivation:** monolayer

**Conditions for cultivation:** medium – DMEM/F12  
serum - FBS 10%

subculture procedure - cells detach from flask using trypsin 0.25%: EDTA 0.02% (1:3), split ratio: 1:3, optimal population density 4.0- 5.0x10<sup>4</sup> cells/cm<sup>2</sup>

cryoconservation - growth medium, 10% DMSO, 1-1,5x10<sup>6</sup> cells/ml in ampule



**Viability after cryoconservation:** 85% (0 passage, dye trypan blue)

**Sterility:** tests for bacteria, fungi and mycoplasma were negative.

**Species:** karyological analysis.

**Karyology:** 2n=46, modal number of chromosomes 46 (98.0%), normal human karyotype (46, XX), number of poliploid cells 1.2%.

**DNA profile (STR):**

Amelogenin:	X,	X
CSF1PO:	10,	12
D13S317:	11,	11
D16S539:	12,	12
D5S818:	7,	11
D7S820:	10,	11
THO1:	6,	7
TPOX:	8,	8
vWA:	15,	16

**Plating efficiency:** 2.4%

**Other properties:** finite lifetime culture; average population doubling time 26.8 h; The presence of surface antigens specific for human MSCs: CD44, CD73, CD90, CD105, HLA-ABC, vimentin and the lack of CD34, HLA-DR; The ability to induced differentiation into osteogenic, chondrogenic and adipogenic directions.

**Applications:** cell biology, biotechnology.

**Collections:** SPBIC.

**Origin:** human, mesenchymal stem cells from jelly of the umbilical cord.

Tsitologiya. 2024. 66 (4):

**Morphology:** fibroblast-like

**Mode of cultivation:** monolayer

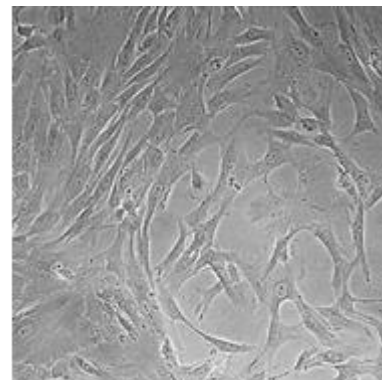
**Conditions for cultivation:**

medium –DMEM/F12

serum - FBS 10%

subculture procedure - cells detach from flask using trypsin 0.25% : EDTA 0.02% (1:3), split ratio: 1:3-1:4, optimal population density 5.0.- 6.0 x10<sup>3</sup> cells/cm<sup>2</sup>

cryoconservation - growth medium, 10%DMSO, 0.5-1.0x10<sup>6</sup> cells/ml in ampule.



**Viability after cryoconservation:** 96% (0 passage, dye trypan blue)

**Sterility:** tests for bacteria, fungi and mycoplasma were negative.

**Species:** karyological analysis.

**Karyology:** 2n=46, modal number of chromosomes 46,XY (98.0 %), normal human karyotype (46, XY), The proportion of polyploidy cells is 2.0, 3.1, 0.6% at passages 6, 16 и 24, respectively.

**DNA profile (STR):**

Amelogenin:	X,	Y
CSF1PO:	10,	11
D13S317:	8,	11
D16S539:	12,	13
D5S818:	12,	13
D7S820:	9,	10
THO1:	8,	9,3
TPOX:	8,	8
vWA:	14,	17

**Other properties:** finite lifetime culture; The stage of active replicative senescence occurs at 24 passage.

The presence of surface antigens specific for human MSCs: CD44, CD73, CD90, CD105, HLA-ABC and the lack of CD34, CD45, HLA-DR.

The ability to induced differentiation into osteogenic, adipogenic and chondrogenic directions; **Applications:** cell biology, biotechnology.

**Collections:** SPBIC.

## NAMALVA

**Origin:** human, Burkitt lymphoma.

Cancer 1969. 23: 64-79; Int.J.Cancer 1972. 10: 44-57; Int.J.Cancer 1973. 12: 396-408; J. Clin. Microbiol. 1975, 1: 116; Antimicrob. Agents Chemother. 1979. 15: 420; Mamaeva S.E. Cell Culture Methods. L., Nauka. 1988: 78-98; Atlas of chromosomes of human and animal cell lines, S.E. Mamaeva, 2002. Moscow, Scientific world

**Morphology:** lymphoblast-like

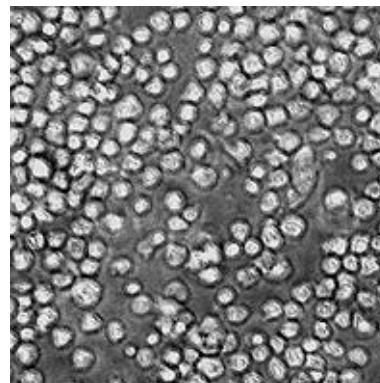
**Mode of cultivation:** suspension

**Conditions for cultivation:** medium - RPMI 1640

serum - FBS 10 %

subculture procedure - optimal population density  $3.0-9.0 \times 10^5$  cells/ml

cryoconservation - growth medium, 5-10% DMSO,  $5.0 \times 10^6$  cells/ml in ampule



**Viability after cryoconservation:** 80% (0 passage, dye trypan blue)

**Sterility:** tests for bacteria, fungi and mycoplasma were negative

**Species:** karyological and isoenzymological (LDH, G6PD) analysis

**Karyology:**  $2n=46$ , variability in the range between 36-48 chromosomes modal number of chromosomes 47, number of markers - 13 (differential dye, G-bandig), number of polyploid cells 2.0%.

**DNA profile (STR):**

Amelogenin:	X,	X
CSF1PO:	10,	12
D13S317:	11,	12
D16S539:	9,	9
D5S818:	12,	13
D7S820:	11,	11
THO1:	7,	9,3
TPOX:	6,	11
vWA:	14,	14

**Other properties:** virus susceptibility: vesicular stomatitis, Sendai.

Secretion of monoclonal antibody (Ig M, lambda light chain).

Support replication of Semliki Forest virus.

**Applications:** biotechnology (interferon  $\alpha$  production), virology, cell biology.

**Collections:** ATCC CRL 1432; ECACC 87060801; DSM (ACC 24); SPBII; MWIIW; SPBIC.



**Origin:** human, kidney carcinoma

Folia Biol. 1988. 34: 308.

**Morphology:** epithelial-like

**Mode of cultivation:** monolayer

**Conditions for cultivation:** medium - DMEM

serum - FBS 10%

subculture procedure - cells detach from flask using trypsin 0.25%: EDTA 0.02% (1:3), split ratio 1:2 - 1:4

cryoconservation - growth medium, 8-10% DMSO,  $1.0 \times 10^6$  cells/ml in ampule

**Viability after cryoconservation:** 90% (0 passage, dye trypan blue)

**Sterility:** tests for bacteria, fungi and mycoplasma were negative

**Species:** karyological and isoenzymological (LDH, G6PD) analysis

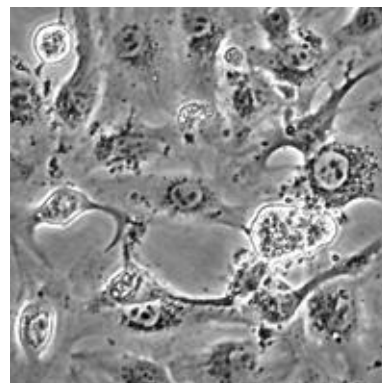
**Karyology:**  $2n=46$ , modal number of chromosomes 75, number of markers - 2 (differential dye)

<b>DNA profile (STR):</b>	Amelogenin:	X,	Y
	CSF1PO:	10,	12
	D13S317:	10,	12
	D16S539:	11,	12
	D5S818:	7,	11
	D7S820:	8,	10
	THO1:	9,	9
	TPOX:	8,	11
	vWA:	16,	18

**Tumorigenicity:** non tumorigenic in nude mice

**Applications:** tumorigenicity, cell biology

**Collections:** SPBIC



**Origin:** human, ovarian teratocarcinoma, ascitic fluid

J.Natl.Cancer Inst. 1974. 52: 921; In Vitro 1974. 10: 382; Int.J.Cancer 1980. 25: 19-32; Atlas of chromosomes of human and animal cell lines, S.E. Mamaeva, 2002. Moscow, Scientific world.

**Morphology:** epithelial-like

**Mode of cultivation:** monolayer

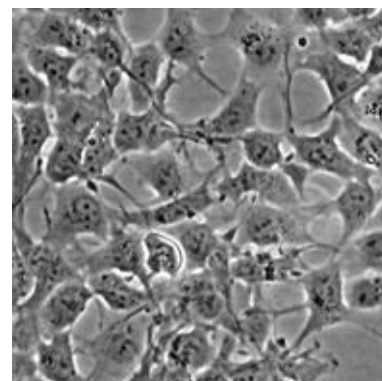
**Conditions for cultivation:** medium - DMEM

serum - FBS 10%

subculture procedure - cells detach from flask using trypsin 0.25%: EDTA 0.02% (1:3), split ratio 1:3 - 1:6, optimal

population density  $1.0-3.0 \times 10^4$  cells/cm<sup>2</sup>

cryoconservation - growth medium, 10% DMSO,  $1.0-2.0 \times 10^6$  cells/ml in ampule



**Viability after cryoconservation:** 87% (0 passage, dye trypan blue)

**Sterility:** tests for bacteria, fungi and mycoplasma were negative

**Species:** karyological and isoenzymological (LDH, G6PD) analysis

**Karyology:**  $2n=46$ , variability in the range between 33-47 chromosomes, modal number of chromosomes 46, pseudodiploid, number of markers - 2 (differential dye), number of polyploid cells 3.0%.

**DNA profile (STR):**

Amelogenin:	X,	X
CSF1PO:	9,	12
D13S317:	9,	10
D16S539:	9,	12
D5S818:	11,	11
D7S820:	9,	9
THO1:	7,	9
TPOX:	11,	11
vWA:	15,	17

**Tumorigenicity:** tumorigenic in nude mice

**Other properties:** chemotaxis, chemoinvasion, matrigel invasion.

**Applications:** tumorigenicity, cell biology.

**Collections:** ATCC CRL 1572; ECACC 90013101; ICLC HTL 97002; SPBIC.

**Origin:** human, pancreatic carcinoma  
Int.J.Cancer 1975. 15: 741-747.

**Morphology:** epithelial-like

**Mode of cultivation:** monolayer

**Conditions for cultivation:** medium - DMEM

serum - FBS 10%

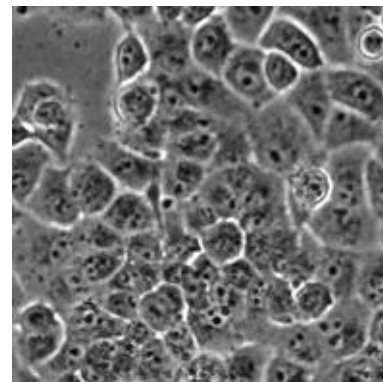
subculture procedure - cells detach from flask using trypsin 0.25%: EDTA 0.02%

(1:1), split ratio 1:2 - 1:4, optimal

population density  $2.0-4.0 \times 10^4$  cells/cm<sup>2</sup>

cryoconservation - growth medium, 10%

DMSO,  $1.0-1.5 \times 10^6$  cells/ml in ampule



**Viability after cryoconservation:** 90% (0 passage, dye trypan blue)

**Sterility:** tests for bacteria, fungi and mycoplasma were negative

**Species:** karyological and isoenzymological (LDH, G6PD) analysis

**Karyology:**  $2n = 46$ , modal number of chromosomes 61 and 63, number of markers - 4 (differential dye), number of polyploid cells 8.5%.

**DNA profile (STR):**

Amelogenin:	X,	X
CSF1PO:	10,	12
D13S317:	11,	11
D16S539:	11,	11
D5S818:	11,	13
D7S820:	8,	10
THO1:	7,	8
TPOX:	8,	11
vWA:	15,	15

**Other properties:** isoenzymes G6PD, B.

**Applications:** tumorigenicity:

**Collections:** ATCC CRL 1469; ECACC 87092802; SPBIC.

**Origin:** human, Burkitt lymphoma

Lancet 1964. 1: 238; J.Bact. 1965. 89: 252; J. Clin. Pathol. 1965. 18: 261; J.Natl.Cancer Inst. 1965. 34: 231; J.Natl.Cancer Inst. 1966. 37: 547; Trans. NY Acad. Sci. 1966. 29: 61; Atlas of chromosomes of human and animal cell lines, S.E. Mamaeva, 2002. Moscow, Scientific world.

**Morphology:** lymphoblast-like

**Mode of cultivation:** suspension

**Conditions for cultivation:** medium - RPMI 1640

serum - FBS 10%

subculture procedure - optimal

population density  $3.0-9.0 \times 10^5$  cells/ml

cryoconservation - growth medium, 10%

DMSO,  $5.0 \times 10^6$  cells/ml in ampule

**Viability after cryoconservation:** 78-88% (0 passage, dye trypan blue)

**Sterility:** tests for bacteria, fungi and mycoplasma were negative

**Species:** karyological and isoenzymological (LDH, G6PD) analysis (LDH, G6PD)

**Karyology:**  $2n=46$ , variability in the range between 43-48 chromosomes, modal number of chromosomes 48, number of markers - 8, number of polyploid cells 4.0%

**DNA profile (STR):**

Amelogenin:	X,	Y
CSF1PO:	10,	12
D13S317:	13,	13
D16S539:	8,	11
D5S818:	10,	13
D7S820:	10,	10
THO1:	6,	7
TPOX:	8,	13
vWA:	16,	19

**Plating efficiency:** 40%

**Other properties:** virus susceptibility: simian retrovirus D, arboviruses.

Isoenzymes G6PD, B.

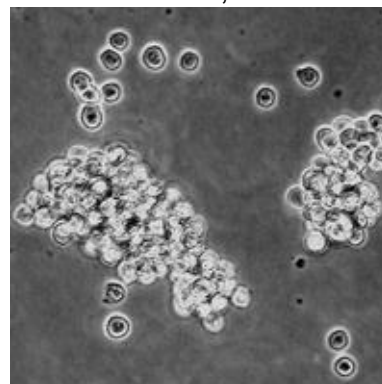
HLA cell line phenotype A (1, 3).

Erythrocyte rosette tests: E, 0; EA, 1%; EAC, 34%.

Positive for EBNA, but does not contain the EBV.

**Applications:** B-cell differentiation, immunology, antitumor testing, virology.

**Collections:** ATCC CCL 86; ECACC 85011429; MWIIW; SPBIC.



**Origin:** human, embryonic rhabdomyosarcoma.

J. Virol. 1967. 1: 326; Cancer 1969. 24: 520-526; Atlas of chromosomes of human and animal cell lines, S.E. Mamaeva, 2002. Moscow, Scientific world.

**Morphology:** spindle-shaped cells and large multinucleated cells.

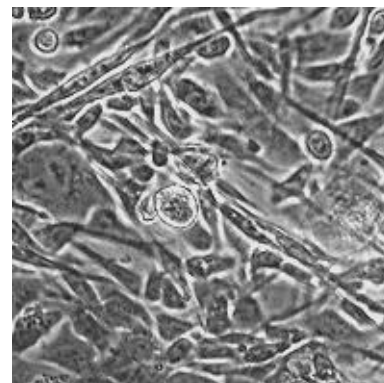
**Mode of cultivation:** monolayer

**Conditions for cultivation:** medium - DMEM

serum - FBS 10%

subculture procedure - cells detach from flask using trypsin 0.25%: EDTA 0.02% (1:2), split ratio 1:3, optimal population density  $4.0 \times 10^4$  cells/cm<sup>2</sup>

cryoconservation - growth medium, 5-10% DMSO,  $1.5-2.0 \times 10^6$  cells/ml in ampule



**Viability after cryoconservation:** 80% (0 passage, dye trypan blue)

**Sterility:** tests for bacteria, fungi and mycoplasma were negative

**Species:** karyological and isoenzymological (LDH, G6PD) analysis

**Karyology:**  $2n=46$ , variability in the range between 45-50 chromosomes, modal number of chromosomes 49, some cells have microchromosomes, number of polyploid cells 3.0%.

**DNA profile (STR):**

Amelogenin:	X,	X
CSF1PO:	10,	11
D13S317:	13,	13
D16S539:	10,	11
D5S818:	11,	11
D7S820:	8,	12
THO1:	9.3,	9.3
TPOX:	9,	9
vWA:	18,	18

**Other properties:** virus susceptibility: poliovirus 1, vesic. stomatitis, herpes simplex, vaccinia, cytomegalovirus, parainfluenza, rotaviruses.

Isoenzymes G6PD, B.

Myoglobin secretion; myoglobin and myosin-ATPase activity.

**Applications:** differentiation, biochemistry, genetics, tumorigenicity, cell biology.

**Collections:** ATCC CCL 136; ECACC 85111502; MWIIW; SPBII; ESCC; SPBIC.

**Origin:** human, leukocytes of peripheral blood from healthy male.

J.Natl.Cancer Inst. 1969. 43: 1119; Atlas of chromosomes of human and animal cell lines, S.E. Mamaeva, 2002. Moscow, Scientific world.

**Morphology:** lymphoblast-like

**Mode of cultivation:** suspension

**Conditions for cultivation:** medium - RPMI 1640

serum - FBS 20%

subculture procedure - optimal

population density  $3.0-4.0 \times 10^5$  cells/ml

cryoconservation - growth medium, 10%

DMSO,  $3.0-4.0 \times 10^6$  cells/ml in ampule

**Viability after cryoconservation:** 75% (0 passage, dye trypan blue)

**Sterility:** tests for bacteria, fungi and mycoplasma were negative

**Species:** karyological and isoenzymological (LDH, G6PD) analysis

**Karyology:**  $2n=46$ , variability in the range between 43-48 chromosomes, modal number of chromosomes 47, number of markers - 1 is in all cells; (differential dye), number of polyploid cells 5.6%

**DNA profile (STR):**

Amelogenin:	X,	Y
CSF1PO:	10,	10
D13S317:	11,	13
D16S539:	10,	13
D5S818:	12,	13
D7S820:	10,	12
THO1:	6,	9.3
TPOX:	8,	9
vWA:	18,	19

**Plating efficiency:** the cells cannot be plated.

**Other properties:** virus susceptibility: poliovirus 1; vesicular stomatitis (Indiana). IgM secretion (lambda light chain).

Isoenzymes G6PD, B.

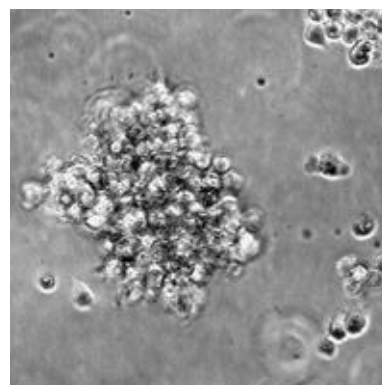
Erythrocyte rosette tests: E, 0; EA, 0; EAC, 19%.

HLA cell line phenotype A2, Aw33, B7, B14.

Positive for EBNA

**Applications:** immunology, biochemistry, cell biology.

**Collections:** ATCC CCL 156; ECACC 85112106; SPBIC.



**Origin:** human, nasal septum carcinoma (Pleural effusion)

Cancer 1964. 17: 170; Exp. Cell Res. 1965. 39: 190; Atlas of chromosomes of human and animal cell lines, S.E. Mamaeva, 2002. Moscow, Scientific world.

**Morphology:** epithelial-like

**Mode of cultivation:** monolayer

**Conditions for cultivation:** medium - EMEM

serum - FBS 10%

other components - NEAA 1%

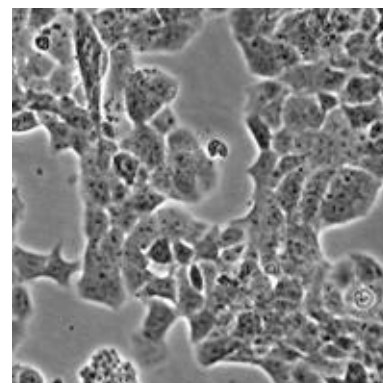
subculture procedure - cells detach from flask using trypsin 0.25%: EDTA 0.02%

(1:3), split ratio 1:2 - 1:3, optimal

population density  $2.0-4.0 \times 10^4$  cells/cm<sup>2</sup>

cryoconservation - growth medium, 10%

DMSO,  $1.0-2.0 \times 10^6$  cells/ml in ampule



**Viability after cryoconservation:** 85% (0 passage, dye trypan blue)

**Sterility:** tests for bacteria, fungi and mycoplasma were negative

**Species:** karyological and immunofluorescent analysis

**Karyology:**  $2n = 46$ , variability in the range between 44-46 chromosomes, modal number of chromosomes 46, pseudodiploid, number of markers - 7 (differential dye), number of polyploid cells 2.2%

**DNA profile (STR):**

Amelogenin:	X,	Y
CSF1PO:	9,	11
D13S317:	11,	12
D16S539:	11,	12
D5S818:	12,	13
D7S820:	8,	11
THO1:	6,	8
TPOX:	8,	8
vWA:	16,	18

**Plating efficiency:** 2%

**Other properties:** virus susceptibility: poliovirus 1, herpes simplex, vesic. stomatitis (Indiana).

Isoenzymes G6PD, B.

Mucopolysaccharide production

**Applications:** tumorigenicity, cell biology.

**Collections:** ATCC CCL 30; ECACC 88031602; SPBIC.

**Origin:** human, myeloma

Proc.Soc.Exp.Biol.Med. 1967, 125: 1246-1250; Atlas of chromosomes of human and animal cell lines, S.E. Mamaeva, 2002. Moscow, Scientific world.

**Morphology:** lymphoblast-like

**Mode of cultivation:** suspension

**Conditions for cultivation:** medium - RPMI 1640

serum - FBS 10%

subculture procedure - optimal  
population density  $5.0 \times 10^5$  cells/ml

cryoconservation - growth medium, 10%  
DMSO,  $5.0 \times 10^6$  cells/ml in ampule

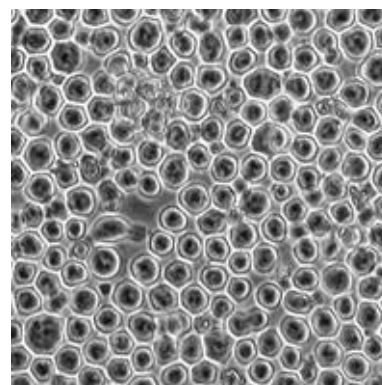
**Viability after cryoconservation:** 55% (0 passage, dye trypan blue)

**Sterility:** bacteria, fungi and mycoplasma were negative

**Species:** karyological and isoenzymological (LDH, G6PD) analysis

**Karyology:**  $2n=46$  variability in the range between 57-73 chromosomes, modal number of chromosomes 67-70, number of markers - 23 (differential dye), number of polyploid cells 10 %

<b>DNA profile (STR):</b>	Amelogenin:	X,	Y
	CSF1PO:	12,	12
	D13S317:	11,	11
	D16S539:	9,	9
	D5S818:	11,	13
	D7S820:	9,	10
	THO1:	8,	8
	TPOX:	8,	11
	vWA:	16,	18



**Plating efficiency:** the cells cannot be plated.

**Other properties:** virus susceptibility: poliovirus 1, vesicular stomatitis (Indiana Strain), herpes simplex, vaccinia.

Isoenzymes G6PD, A.

Secrete  $\lambda$ -type light chains of immunoglobulin.

Erythrocyte rosette tests: E, 0; EA, 1%; EAC, 13%.

HLA cell line phenotype : Aw 19, B 15, B 37, Cw 2.

**Applications:** cell biology, tumorigenicity: , immunology, biotechnology (production Ig)

**Collections:** ATCC CCL 155, ECACC 87012702; SPBIC.



**Origin:** embryonic stem cells (ESC) from blastocyst 5-6 days  
 Science. 1998. 282: 1145 – 1147; Ontogenez. 2011. 42 (4): 249 – 263; Tsitologiya. 2012. 54 (1): 5 – 16.

**Morphology:** colonies of rounded cells

**Mode of cultivation:** monolayer; colonies attached to the feeder layer of mitotically inactivated (mitomycin C) cells of line FetMSC

**Conditions for cultivation:** medium – Knockout DMEM

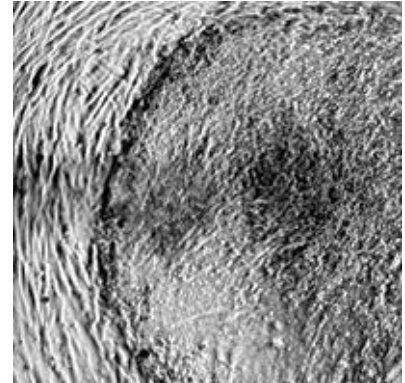
serum - Knockout serum replacement

other components - NEAA 1%, L-glutamine 2mM, 2- mercaptoethanol 0.1 mM, bFGF – 8ng/ml

subculture procedure - mechanical reseeded of culture ESC carried out

under the control of the microscope by cutting the colony into fragments using a single scalpel and transfer them onto a new layer feeder; daily changing growth medium; subculture every 5-6 days

cryoconservation - growth medium, 10% DMSO,  $5 \times 10^5$  cells/ml in ampule



**Viability after cryoconservation:** 60% (0 passage, dye trypan blue)

**Sterility:** tests for bacteria, fungi and mycoplasma were negative.

**Species:** karyological analysis.

**Karyology:**  $2n=46$ , modal number of chromosomes 46 (98.0%), normal human karyotype (46, XX), number of poliploid cells 0.2%.

**DNA profile (STR):**

Amelogenin:	X,	X
CSF1PO:	12,	13
D13S317:	8,	11
D16S539:	9,	12
D5S818:	9,	11
D7S820:	8,	10, 12
THO1:	6,	9.3
TPOX:	10,	11
vWA:	17,	17

**Other properties:** immortalized line; passed through more than 120 cell population doublings; average population doubling time 28.2 h; The presence of surface antigens specific for human ESC: SSEA-4, TRA-1-60, Oct-4, Nanog; The ability to differentiation into the derivatives of the 3 germ layers and forming teratomas, containing these derivatives.

**Applications:** cell biology, embryology, biotechnology.

**Collections:** SPBIC.

**Origin:** mesenchymal stem cells from human embryonic stem cells (ESC).

Tsitologiya. 2012. 54 (1): 5 – 16; Tissue Eng Part A. 2010. 16:705 – 715.

**Morphology:** fibroblast-like

**Mode of cultivation:** monolayer

**Conditions for cultivation:** medium –  $\alpha$ -MEM

serum - FBS 10%

subculture procedure - cells detach from flask using trypsin 0.25%: EDTA 0.02% (1:3), split ratio: 1:4, optimal population density 4.0- 5.0x10<sup>4</sup> cells/cm<sup>2</sup>

cryoconservation - growth medium, 10% DMSO, 1.5-2.0x10<sup>6</sup> cells/ml in ampule

**Viability after cryoconservation:** 90% (0 passage, dye trypan blue)

**Sterility:** tests for bacteria, fungi and mycoplasma were negative.

**Species:** karyological analysis.

**Karyology:** 2n=46, modal number of chromosomes 46 (100.0±1.0%), normal human karyotype (46, XX), number of poliploid cells 0.9%.

**DNA profile (STR):**

Amelogenin:	X,	X
CSF1PO:	12,	13
D13S317:	8,	11
D16S539:	9,	12
D5S818:	9,	11
D7S820:	10,	12
THO1:	6,	9.3
TPOX:	10,	11
vWA:	17,	17

**Other properties:** finite lifetime culture; average population doubling time 25.5 h; The presence of surface antigens specific for human MSCs: CD44, CD73, CD90, CD105, HLA-ABC, and the lack of CD34, HLA-DR; The ability to induced differentiation into osteogenic, chondrogenic and adipogenic directions.

**Applications:** cell biology, biotechnology, feeder for cultivation embryonic stem cells.

**Collections:** SPBIC.

**Origin:** human, mesenchymal stem cells from embryonic stem cells of line SC7.

**Tsitologiya.** 2022. 64 (5): 713 – 727.

**Morphology:** fibroblast-like

**Mode of cultivation:** monolayer

**Conditions for cultivation:** medium – DMEM/F12

serum - FBS 10%

subculture procedure - cells detach from flask using 0.25% solution trypsin:EDTA, split ratio: 1:3, optimal population density  $0,7-2.0 \times 10^4$  cells/cm<sup>2</sup>

cryoconservation - growth medium, 10% DMSO,  $2.5 \times 10^6$  cells/ml in ampule.



**Viability after cryoconservation:** 80% (0 passage, dye trypan blue)

**Sterility:** tests for bacteria, fungi and mycoplasma were negative.

**Species:** karyological analysis.

**Karyology:**  $2n=46$ , modal number of chromosomes 46 ( $99.0 \pm 1.0$  %), normal human karyotype (46, XY), number of poliploid cells  $5.7 \pm 0.9$ %.

**DNA profile (STR):**

Amelogenin:	X,	Y
CSF1PO:	10,	12
D13S317:	8,	11
D16S539:	11,	11
D5S818:	13,	13
D7S820:	9,	10
THO1:	6,	6
TPOX:	8,	11
vWA:	17,	18

**Plating efficiency:** 0.0 %.

**Other properties:** finite lifetime culture; at the 6 passage the average time of one doubling of the cell population is  $36.0 \pm 0.5$  h. The time of the active (logarithmic) growth phase is 96 h. The stage of active replicative senescence begins at passage 13; cell death is observed at the 20th passage. The presence of surface antigens specific for human MSCs: CD44, CD73, CD90, CD105, vimentin, HLA-ABC and the lack of CD34, CD45, HLA-DR; directed differentiation in the osteogenic, adipogenic and chondrogenic directions at the 6th passage; the absence of adipogenic and osteogenic differentiation at the stage of active replicative senescence (P.13).

**Applications:** cell biology, biotechnology.

**Collections:** SPBIC.

## SK-HEP-1

**Origin:** human, liver adenocarcinoma (ascitic fluid).

J.Natl.Cancer Inst. 1977. 58: 209; J.Natl.Cancer Inst. 1977. 59: 221.

**Morphology:** epithelial-like

**Mode of cultivation:** monolayer

**Conditions for cultivation:** medium - EMEM

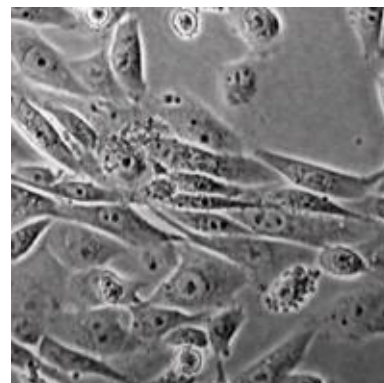
serum - FBS 10%

other components - NEAA 1%, sodium pyruvate 1mM.

subculture procedure - cells detach from flask using trypsin 0.25%: EDTA 0.02% (1:3), split ratio 1:2 - 1:4, optimal

population density  $2.0-4.0 \times 10^4$  cells/cm<sup>2</sup>

cryoconservation - growth medium, 5%DMSO,  $1.0 \times 10^6$  cells/ml in ampule



**Viability after cryoconservation:** 93% (0 passage, dye trypan blue)

**Sterility:** tests for bacteria, fungi and mycoplasma were negative

**Species:** karyological analysis

**Karyology:**  $2n=46$ , variability in the range between 58-64 chromosomes, modal number of chromosomes 60-61, number of markers - 8 (differential dye), 50% of cells have large acrocentric chromosome, number of polyploid cells 0.4%.

**DNA profile (STR):**

Amelogenin:	X,	X
CSF1PO:	11,	12
D13S317:	8,	12
D16S539:	12,	12
D5S818:	10,	13
D7S820:	8,	11
THO1:	7,	9
TPOX:	9,	9
vWA:	14,	17

**Tumorigenicity:** tumorigenic in nude mice

**Other properties:** isoenzymes Me-2, 1-2; PGM<sub>3</sub>,1; PGM<sub>1</sub>, 2; ES D,1; AK 1,1; GLO-1,1; G6PD,B.

bFGF production.

**Applications:** tumorigenicity:

**Collections:** ATCC HTB 52; ECACC 91091816; SPBIC.

**Origin:** human, neuroblastoma (metastasis to supra-orbital area)  
Cancer Res. 1973. 33: 2643; In Vitro 1973. 8: 410; Cancer Res. 1977. 37: 1364; Atlas of chromosomes of human and animal cell lines, S.E. Mamaeva, 2002. Moscow, Scientific world.

**Morphology:** epithelial-like and neuroblast-like

**Mode of cultivation:** monolayer

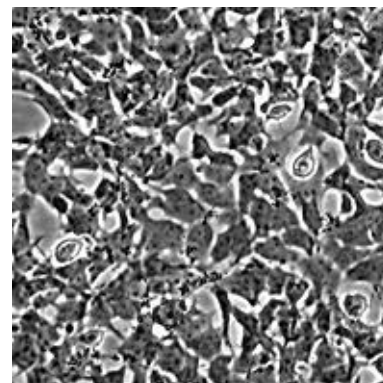
**Conditions for cultivation:** medium - EMEM

serum - FBS 10%

other components - NEAA 1%

subculture procedure - cells detach from flask using trypsin 0.25%: EDTA 0.02% (1:3), split ratio 1:2 - 1:5

cryoconservation - growth medium, 5-10% DMSO, 1.0-1.5x10<sup>6</sup> cells/ml in ampule



**Viability after cryoconservation:** 90% (0 passage, dye trypan blue)

**Sterility:** tests for bacteria, fungi and mycoplasma were negative

**Species:** karyological analysis

**Karyology:** 2n= 46, variability in the range between 44-47 chromosomes, modal number of chromosomes 46, pseudodiploid, number of markers - 15 (differential dye), number of polyploid cells 1.2%

**DNA profile (STR):**

Amelogenin:	X,	X
CSF1PO:	10,	10
D13S317:	11,	11
D16S539:	12,	12
D5S818:	11,	11
D7S820:	8,	8
THO1:	9.3,	9.3
TPOX:	9,	11
vWA:	17,	18

**Tumorigenicity:** tumorigenic: produce neuroblastoma in nude mice; produce tumors in the cheek pouch of the hamster.

**Other properties:** isoenzymes Me-2,2; PGM<sub>3</sub>,1-2; PGM<sub>1</sub>,1; ES D,2; AK-1,1; GLO-1,1-2; G6PD,B.

Catecholamine production.

**Applications:** neurophysiology, biochemistry.

**Collections:** ATCC HTB 10; SPBIC.

**Origin:** human, uterine leiomyosarcoma.

J. Natl.Cancer Inst. 1977. 59: 221-226; Cancer Genet. Cytogenet. 1988, 33: 77-81

**Morphology:** epithelial-like

**Mode of cultivation:** monolayer (weak adhesion)

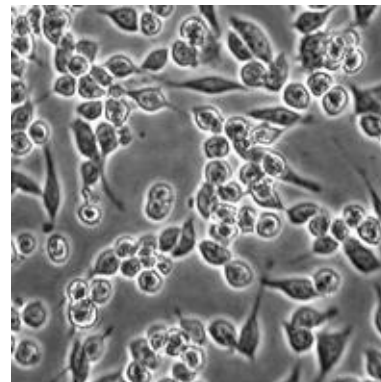
**Conditions for cultivation:** medium - EMEM

serum - FBS 10%

other components - NEAA 1%

subculture procedure - cells detach from flask using trypsin 0.25%: EDTA 0.02% (1:3), split ratio 1.3 -1:5

cryoconservation - growth medium, 8%DMSO,  $1.0-2.0 \times 10^6$  cells/ml in ampule



**Viability after cryoconservation:** 82% (0 passage, dye trypan blue)

**Sterility:** tests for bacteria, fungi and mycoplasma were negative

**Species:** karyological analysis

**Karyology:**  $2n = 46$ , variability in the range between 44-48 chromosomes, modal number of chromosomes 46, normal human karyotype (46, XX), number of polyploid cells 0.6%.

**DNA profile (STR):**

Amelogenin:	X,	X
CSF1PO:	10,	11
D13S317:	10,	13
D16S539:	12,	14
D5S818:	10,	11
D7S820:	9,	10
THO1:	7,	7
TPOX:	8,	8
vWA:	16,	16

**Tumorigenicity:** tumorigenic in nude mice

**Other properties:** isoenzymes Me2,1-2; PGM<sub>3</sub>,1; PGM<sub>1</sub>,1; ESD,1; AK 1,1; GLO-1,1-2; G6PD,B

**Applications:** tumorigenicity, cytogenetics, cell biology.

**Collections:** ATCC HTB 115; SPBIC.

**Origin:** human, rectum adenocarcinoma.

Cancer Res. 1976. 36: 4562- 4569; Cytology ( Russ.) 1992. 34: 63-64.; Atlas of chromosomes of human and animal cell lines, S.E. Mamaeva, 2002. Moscow, Scientific world.

**Morphology:** epithelial-like

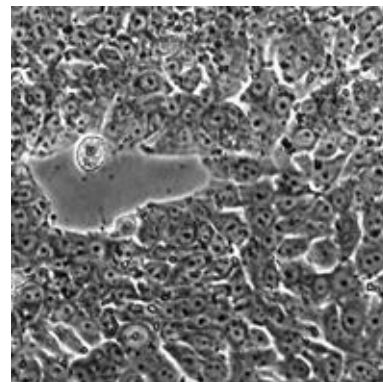
**Mode of cultivation:** monolayer

**Conditions for cultivation:** medium - L-15 (Leibovitz)

serum - FBS 10%

subculture procedure - cells detach from flask using trypsin 0.25%: EDTA 0.02% (1:3), split ratio 1:2 (subcultivation in 14-18 days), optimal population density 2.0-4.0x10<sup>4</sup> cells/cm<sup>2</sup>

cryoconservation - growth medium, 10% DMSO, 2.0-3.0x10<sup>6</sup> cells/ml in ampule



**Viability after cryoconservation:** 80% (0 passage, dye trypan blue)

**Sterility:** tests for bacteria, fungi and mycoplasma were negative

**Species:** karyological analysis

**Karyology:** 2n= 46, variability in the range between 34-41 chromosomes, modal number of chromosomes 40, number of markers - 11 (differential dye), number of polyploid cells 10%.

**DNA profile (STR):**

Amelogenin:	X,	X
CSF1PO:	10,	10
D13S317:	13,	13
D16S539:	12,	12
D5S818:	12,	12
D7S820:	9,	12
THO1:	9.3,	9.3
TPOX:	8,	9
vWA:	15,	16

**Plating efficiency:** 2%.

**Tumorigenicity:** tumorigenic in nude mice.

**Other properties:** isoenzymes G6PD, B; PGM<sub>3</sub>, 1; PGM<sub>1</sub>, 1; PGD, A; ES D, 1. CEA production.

**Applications:** tumorigenicity, cell biology.

**Collections:** ATCC CCL 235; ECACC 91031104; SPBIC.

**Origin:** human, bladder carcinoma.

Int. J. Cancer 1970.5: 310; Int. J.Cancer 1971. 8: 503; Int. J. Cancer 1973.11: 765; Tissue Antigens. 1978.11:279.

**Morphology:** epithelial-like

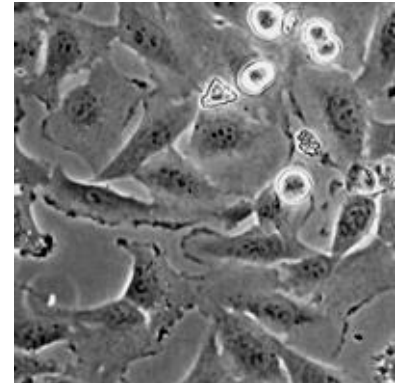
**Mode of cultivation:** monolayer

**Conditions for cultivation:** medium - EMEM

serum - FBS 10%

subculture procedure - cells detach from flask using trypsin 0.25%: EDTA 0.02% (1:3), split ratio 1:3-1:5, optimal population density  $1.0 \times 10^5$  cells/cm<sup>2</sup>

cryoconservation - growth medium, 10% DMSO,  $1.0-2.0 \times 10^6$  cells/ml in ampule



**Viability after cryoconservation:** 86% (0 passage, dye trypan blue)

**terility:** tests for bacteria, fungi and mycoplasma were negative

**Species:** karyological and isoenzymological (LDH, G6PD) analysis

**Karyology:**  $2n=46$ , variability in the range between 77-99 chromosomes, modal number of chromosomes 93 without markers (routine, differential dye, C-banding), there are microchromosomes, number of polyploid cells 2.0%.

**DNA profile (STR):**

Amelogenin:	X,	X
CSF1PO:	10,	12
D13S317:	12,	12
D16S539:	9,	9
D5S818:	10,	12
D7S820:	10,	11
THO1:	6,	6
TPOX:	8,	11
vWA:	17,	17

**Tumorigenicity:** tumorigenic

**Other properties:** isoenzymes G6PD,B; Me-2,2-1; PGM 3,1; FUC,2-1; PGM 1,1; ESD,1; ADA,1.

HLA cell line phenotype A (1,3); B (8,18); C (w2, w6), Ek-2, DRw2, w4

**Applications:** virology, tumorigenicity.

**Collections:** ATCC HTB 4; MWIIW, SPBIC



**Origin:** human, glioblastoma.

J. Cell Physiol. 1979. 99: 43-54.

**Morphology:** epithelial-like

**Mode of cultivation:** monolayer

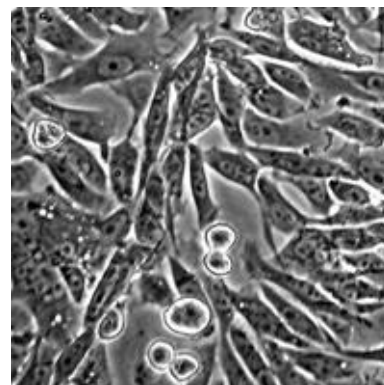
**Conditions for cultivation:** medium - EMEM

serum - FBS 10%

other components - NEAA 1%

subculture procedure - cells detach from flask using trypsin 0.25%: EDTA 0.02% (1:3), split ratio 1:3 -1:6

cryoconservation - growth medium, 10% DMSO,  $1.0 \times 10^6$  cells/ml in ampule



**Viability after cryoconservation:** 90% (0 passage, dye trypan blue)

**Sterility:** tests for bacteria, fungi and mycoplasma were negative

**Species:** karyological analysis

**Karyology:**  $2n = 46$ , modal number of chromosomes 128-132, number of markers - 14-16 (differential dye), number of polyploid cells 1.3%.

**DNA profile (STR):**

Amelogenin:	X,	Y
CSF1PO:	10,	12
D13S317:	13,	13
D16S539:	13,	13
D5S818:	10,	12
D7S820:	9,	10
THO1:	7,	9.3
TPOX:	8,	8
vWA:	17,	20

**Applications:** studies on the mechanisms for cessation of proliferation, cell synchronisation in  $G_1$  phase and ageing.

**Collections:** ATCC CRL 1690; SPBIC.

**Origin:** human, peripheral blood, acute monocytic leukemia from 1-year-old male  
 Int. J. Cancer 1980. 26: 171 – 176; Cancer Res. 1982. 42: 1530; J. Immunol. 1983. 131: 1882; Genes Chromosomes Cancer. 2000. 29: 333 – 338; Atlas of chromosomes of human and animal cell lines, S.E. Mamaeva, 2002. Moscow, Scientific world.

**Morphology:** monocyto-like

**Mode of cultivation:** suspension

**Conditions for cultivation:** medium - RPMI 1640

serum - FBS 10%

other components - 2-mercaptoetanol  
 $2 \times 10^{-5} \text{M}$

subculture procedure - optimal  
 population density  $1.0 - 5.0 \times 10^5$  cells/ml

cryoconservation - growth medium, 10%  
 DMSO,  $4.0 - 6.0 \times 10^6$  cells/ml in ampule

**Viability after cryoconservation:** 80% (0 passage, dye trypan blue)

**Sterility:** tests for bacteria, fungi and mycoplasma were negative

**Species:** karyological analysis

**Karyology:**  $2n = 46$ , modal number of chromosomes 50, number of markers - 8 (differential dye), number of polyploid cells 2.5%.

**DNA profile (STR):**

Amelogenin:	X,	Y
CSF1PO:	11,	13
D13S317:	13,	13
D16S539:	11,	12
D5S818:	11,	12
D7S820:	10,	10
THO1:	5,	8, 9.3
TPOX:	8,	11
vWA:	16,	16

**Other properties:** presence Fc and C3b receptors.

Lack surface and cytoplasmic immunoglobulins.

Produce lysozymes, phagocytic activity.

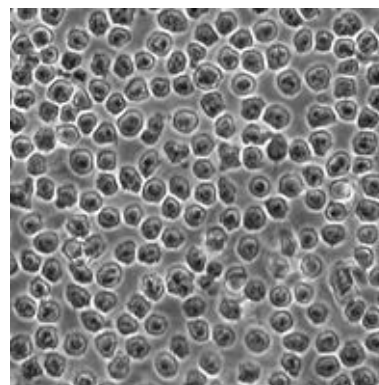
Differentiation into macrophage-like cells.

Induce by herbology ether of monocytic differentiation.

HLA cell phenotype – A2, A9, B5, DRw1, DRw2.

**Applications:** immunology, differentiation, tumorigenicity.

**Collections:** ATCC TIB-202; ECACC 88081201; DSM ACC 16; SPBIC.



**Origin:** human, osteosarcoma.

Int.J.Cancer 1967. 2: 434-447.

**Morphology:** epithelial-like

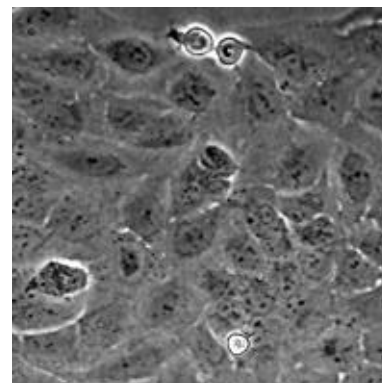
**Mode of cultivation:** monolayer

**Conditions for cultivation:** medium - RPMI 1640

serum - FBS 10

subculture procedure - cells detach from flask using trypsin 0.25%: EDTA 0.02% (1:3), split ratio 1:2 - 1:4

cryoconservation - growth medium, 5-10% DMSO,  $1.0-2.0 \times 10^6$  cells/ml in ampule



**Viability after cryoconservation:** 92% (0 passage, dye trypan blue)

**Sterility:** tests for bacteria, fungi and mycoplasma were negative

**Species:** karyological and isoenzymological (LDH, G6PD) analysis

**Karyology:**  $2n = 46$ , variability in the range between 67-80 chromosomes, modal number of chromosomes 76 and 78-79, number of markers - 22 (differential dye).

**DNA profile (STR):**

Amelogenin:	X,	X
CSF1PO:	12,	13
D13S317:	13,	13
D16S539:	11,	12
D5S818:	8,	11
D7S820:	11,	12
THO1:	6,	6
TPOX:	11,	12
vWA:	14,	18

**Other properties:** isoenzymes PGM1, 1; PGM3, 2; ES D, 1; AK 1, 1; GLO-1, 2; G6PD, B.

**Applications:** tumorigenicity, cell biology.

**Collections:** ATCC HTB 96; SPBIC.

**Origin:** human, histiocytic lymphoma (pleural effusion)

Int.J.Cancer 1976. 17: 565-577; J.Exp.Med. 1976. 143: 1528-1533; J.Immunol. 1980. 125: 463-465; Nature 1979. 279: 328-331; Atlas of chromosomes of human and animal cell lines, S.E. Mamaeva, 2002. Moscow, Scientific world.

**Morphology:** histiomonocitoid

**Mode of cultivation:** suspension

**Conditions for cultivation:** medium - RPMI 1640

serum - FBS 10%

subculture procedure - optimal population density  $2.0-9.0 \times 10^5$  cells/ml

cryoconservation - growth medium, 8-9%DMSO,  $5.0 \times 10^6$  cells/ml in ampule

**Viability after cryoconservation:** 80% (0 passage, dye trypan blue)

**Sterility:** tests for bacteria, fungi and mycoplasma were negative

**Species:** karyological and isoenzymological (LDH, G6PD) analysis

**Karyology:**  $2n=46$ , variability in the range between 59-65 chromosomes, modal number of chromosomes 61, number of markers - 21 (differential dye), number of polyploid cells 3.0%.

**DNA profile (STR):**

Amelogenin:	X,	X
CSF1PO:	10,	12
D13S317:	10,	12
D16S539:	12,	12
D5S818:	10,	12, 13
D7S820:	9,	11
THO1:	6,	9.3
TPOX:	8,	11
vWA:	14,	15

**Other properties:** virus susceptibility: HIV-1, HIV-2, herpes type 6.

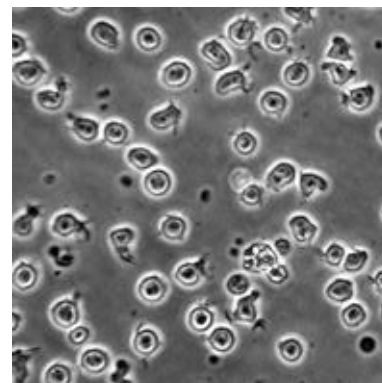
IL-1 production.

Fc and C3 receptors.

Phagocytose antibody-coated erythrocytes and latex beads.

**Applications:** differentiation, virology, cell biology, tumorigenicity.

**Collections:** ATCC CRL 1593; DSM ACC 5; ECACC 85011440; 87010802; ICLC HTL 94002; SPBII; SPBIC.



## WI-38 VA 13 subline 2RA

**Origin:** human, embryonic lung, an SV 40 virus-transformed derivative of the WI-38 cell line.

Ann.Med.Exp.Biol.Fenn.1966. 44:242; J.Natl.Cancer Inst.1964. 32: 917.

**Morphology:** epithelial-like

**Mode of cultivation:** monolayer

**Conditions for cultivation:** medium - EMEM

serum - FBS 10%

other components - NEAA 1%

subculture procedure - cells detach from

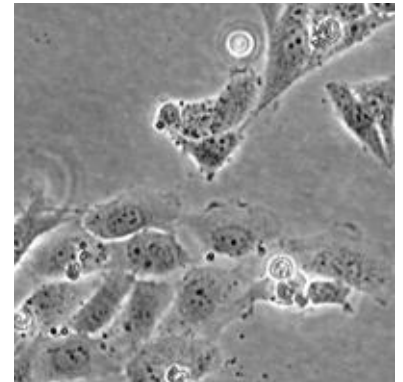
flask using trypsin 0.25%: EDTA 0.02%

(1:3), split ratio 1:2 - 1:4, optimal population

density  $1.0-3.0 \times 10^4$  cells/cm<sup>2</sup>

cryoconservation - growth medium,

5%DMSO,  $1.0 \times 10^6$  cells/ml in ampule



**Viability after cryoconservation:** 80-85% (0 passage, dye trypan blue)

**Sterility:** tests for bacteria, fungi and mycoplasma were negative

**Species:** karyological analysis

**Karyology:**  $2n=46$ , variability in the range between 45-89 chromosomes, modal number of chromosomes 73-78, number of markers - 2-3 (routine dye), 1-6 microchromosomes.

**DNA profile (STR):**

Amelogenin:	X,	X
CSF1PO:	10,	12
D13S317:	11,	11
D16S539:	11,	12
D5S818:	10,	10
D7S820:	9,	11
THO1:	9.3,	9.3
TPOX:	8,	8
vWA:	19,	20

**Plating efficiency:** 15%

**Other properties:** virus susceptibility: herpes simplex, vesicular stomatitis (Indiana), poliovirus 2, reovirus 3.

Isoenzymes G6PD.

Contains SV 40 neo (T) and transplantation antigens.

**Applications:** biochemistry, transformation, virology.

**Collections:** ATCC CCL 75.1; ECACC 85062512; SPBIC.

**Origin:** human, SV 40 virus-transformed fibroblasts from xeroderma pigmentosum patients.

Mol.Cell Biol. 1987. 7: 3353-3357.

**Morphology:** fibroblast-like

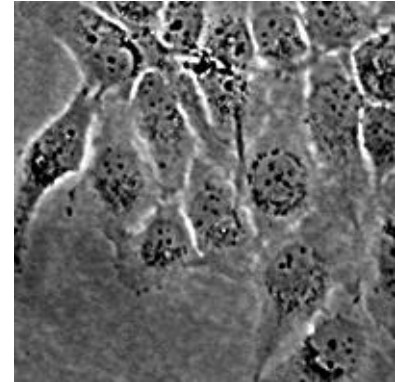
**Mode of cultivation:** monolayer

**Conditions for cultivation:** medium - DMEM

serum - FBS 10%

subculture procedure - cells detach from flask using trypsin 0.25%: EDTA 0.02% (1:3), split ratio 1:3-1:5

cryoconservation - growth medium, 5-8%DMSO,  $1.0 \times 10^6$  cells/ml in ampule



**Viability after cryoconservation:** 90% (0 passage, dye trypan blue)

**Sterility:** tests for bacteria, fungi and mycoplasma were negative

**Species:** karyological and isoenzymological (LDH, G6PD) analysis

**Karyology:**  $2n = 46$ , variability in the range between 55-75 chromosomes, modal number of chromosomes 68-70, number of markers – 19% dicentrics (routine dye), 7% of cells have microchromosomes, number of polyploid cells 5.0%.

**DNA profile (STR):**

Amelogenin:	X,	X
CSF1PO:	12,	12
D13S317:	12,	12
D16S539:	9,	11
D5S818:	11,	12
D7S820:	12,	12
THO1:	9,	9
TPOX:	8,	11
vWA:	17,	17

**Applications:** genetics, tumorigenicity, cell biology.

**Collections:** SPBIC

**Origin:** human, mammary gland carcinoma (ascitic effusion)

Cancer Res. 1978. 38: 3352-3364 и 4327-4339.

**Morphology:** epithelial-like

**Mode of cultivation:** monolayer

**Conditions for cultivation:** medium - RPMI 1640

serum - FBS 10%

subculture procedure - cells detach from

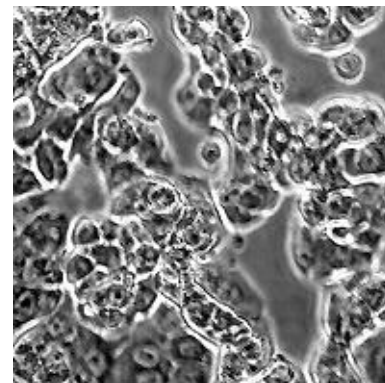
flask using trypsin 0.25%: EDTA 0.02%

(1:3), split ratio 1:2 - 1:3, optimal

population density  $2.0-4.0 \times 10^4$  cells/cm<sup>2</sup>

cryoconservation - growth medium, 10%

DMSO,  $2.0 \times 10^6$  cells/ml in ampule



**Viability after cryoconservation:** 90% (0 passage, dye trypan blue)

**Sterility:** tests for bacteria, fungi and mycoplasma were negative

**Species:** karyological and isoenzymological (LDH, G6PD) analysis

**Karyology:**  $2n=46$ , variability in the range between 55-77 chromosomes, modal number of chromosomes 72, number of markers - 18 (differential dye), number of polyploid cells 0.8%.

**DNA profile (STR):**

Amelogenin:	X,	X
CSF1PO:	10,	11
D13S317:	9,	9
D16S539:	11,	11
D5S818:	13,	13
D7S820:	10,	11
THO1:	7,	9.3
TPOX:	8,	8
vWA:	16,	18

**Other properties:** receptors for estrogen and other steroid hormones.

**Applications:** tumorigenicity, cell biology.

**Collections:** ATCC CRL 1500; ECACC 87012601; SPBIC.

## ANIMAL CELL LINES

35

**Origin:** rat, glioma induced by ethylnitrozourea.

Submitted from Research Institute of Neurosurgery of the Ukrainian Ministry of Health, Kiev; Atlas of chromosomes of human and animal cell lines, S.E. Mamaeva, 2002.

Moscow, Scientific world

**Morphology:** glial

**Mode of cultivation:** monolayer

**Conditions for cultivation:** medium - EMEM

serum - FBS 10%

subculture procedure - cells detach from flask using trypsin 0.25%: EDTA 0.02% (1:3), split ratio 1:5.

cryoconservation - growth medium, 10% DMSO,  $1.0 \times 10^6$  cells/ml in ampule

**Viability after cryoconservation:** 96% (0 passage, dye trypan blue)

**Sterility:** tests for bacteria, fungi and mycoplasma were negative

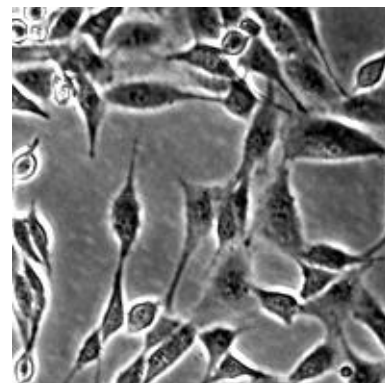
**Species:** karyological and isoenzymological (LDH, G6PD) analysis

**Karyology:**  $2n=42$ , variability in the range between 41-44 (first clone of cell population), 81-86 (second clone of cell population) chromosomes: modal number of chromosomes 82-84, number of markers - 3 (differential dye).

**Tumorigenicity:** tumorigenic

**Applications:** neurobiology, tumorigenicity.

**Collections:** SPBIC





**Origin:** rat, glioma induced by N-methylnitrozourea.  
Exp.Oncol. (Russ) 1982. 2: 27.

**Morphology:** glial

**Mode of cultivation:** monolayer

**Conditions for cultivation:** medium - EMEM

serum - FBS 10%

subculture procedure - cells detach from flask using trypsin 0.25%: EDTA 0.02% (1:3), split ratio 1:5 - 1:8.

cryoconservation - growth medium, 10% DMSO,  $1.0 \times 10^6$  cells/ml in ampule

**Viability after cryoconservation:** 97% (0 passage, dye trypan blue)

**Sterility:** tests for bacteria, fungi and mycoplasma were negative

**Species:** karyological and isoenzymological (LDH, G6PD) analysis

**Karyology:**  $2n = 42$ , variability in the range between 78-85 chromosomes, modal number of chromosomes 81-83, number of markers - 20 (differential dye).

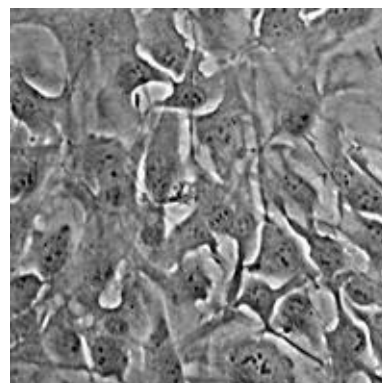
**Plating efficiency:** 45%

**Tumorigenicity:** tumorigenic in syngeneic animals

**Other properties:** secretion of protein S-100

**Applications:** neurobiology, tumorigenicity.

**Collections:** SPBIC



### 3T3B-SV40

**Origin:** mouse BALB/c, embryo, BALB/3T3 clone A31 transformed by SV40.

**Morphology:** fibroblast-like

**Mode of cultivation:** monolayer

**Conditions for cultivation:** medium - DMEM

serum - FBS 10%

subculture procedure - cells detach from flask using trypsin 0.25%: EDTA 0.02% (1:3), split ratio 1:3 - 1:6, optimal population density  $2.0-4.0 \times 10^4$  cells/cm<sup>2</sup>

cryoconservation - growth medium, 10% DMSO,  $2.0 \times 10^6$  cells/ml in ampule

**Viability after cryoconservation:** 80% (0 passage, dye trypan blue)

**Sterility:** tests for bacteria, fungi and mycoplasma were negative

**Species:** karyological, and isoenzymological (LDH, G6PD) and immunofluorescent analysis

**Karyology:**  $2n=40$ , variability in the range between 66-73 chromosomes, modal number of chromosomes 70, 1-2 microchromosomes in 40% of cells.

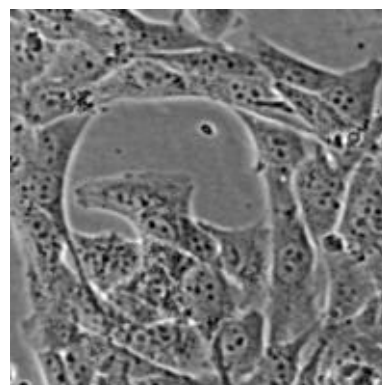
**Plating efficiency:** 30%

**Tumorigenicity:** non tumorigenic

**Other properties:** T antigen in nucleus

**Applications:** tumorigenicity, virology, cell biology.

**Collections:** SPBIC



**Origin:** mouse, embryo, 3T3 Swiss cells transformed by SV 40  
Submitted from «Flow Labs» 1986.

**Morphology:** fibroblast-like

**Mode of cultivation:** monolayer

**Conditions for cultivation:** medium - DMEM

serum - FBS 10%

subculture procedure - cells detach from flask using trypsin 0.25%: EDTA 0.02% (1:3), split ratio 1:3 - 1:5

cryoconservation - growth medium, 10% DMSO,  $1.0 \times 10^6$  cells/ml in ampule

**Viability after cryoconservation:** 90% (0 passage, dye trypan blue)

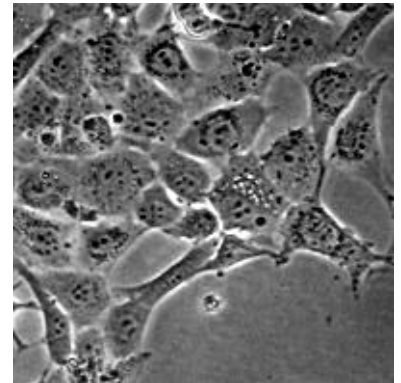
**Sterility:** tests for bacteria, fungi and mycoplasma were negative

**Species:** karyological and isoenzymological (LDH, G6PD) analysis

**Karyology:**  $2n=40$ , variability in the range between 56-72 chromosomes, some cells have large submetacentric and metacentric chromosomes and middle acrocentric chromosome with secondary constriction (routine dye), number of polyploid cells 0.8%.

**Applications:** cell biology

**Collections:** SPBIC



### 3T3 Swiss albino

**Origin:** Swiss mouse, embryo.

J. Cell Biol. 1963. 17: 299-313.

**Morphology:** fibroblast-like

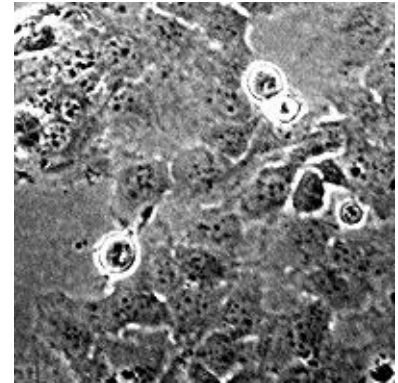
**Mode of cultivation:** monolayer

**Conditions for cultivation:** medium - DMEM

serum - FBS 10%

subculture procedure - cells detach from flask using trypsin 0.25%: EDTA 0.02% (1:1), split ratio 1:3 - 1:6, optimal population density  $5.0 \times 10^3$  -  $1.0 \times 10^4$  cells/cm<sup>2</sup>

cryoconservation - growth medium, 5 – 10% DMSO,  $1.0 \times 10^6$  cells/ml in ampule



**Viability after cryoconservation:** 90% (0 passage, dye trypan blue)

**Sterility:** tests for bacteria, fungi and mycoplasma were negative

**Species:** karyological and isoenzymological (LDH, G6PD) analysis

**Karyology:**  $2n=40$ , variability in the range between 65-73 chromosomes, modal number of chromosomes 69-71, number of markers - 2-3, small acrocentric chromosomes (routine dye), some cells have 1-2 microchromosomes, number of polyploid cells 1.2%.

**Plating efficiency:** 20 %.

**Tumorigenicity:** non tumorigenic

**Other properties:** virus susceptibility: herpes simplex, Sendai, vesic. stomatitis (Indiana), vaccinia.

Contact inhibition of growth.

**Applications:** biochemistry, differentiation, virology, genetical transformation, tumorigenicity.

**Collections:** ATCC CCL 92; ECACC 85022108; SPBIC.

**Origin:** Swiss mouse, embryo.

Keratinocyte methods by I. and F. Walt. Cambridge University Press 1994. P.5-12.

**Morphology:** fibroblast-like

**Mode of cultivation:** monolayer

**Conditions for cultivation:** medium - DMEM

serum - FBS 10%

subculture procedure - cells detach from flask using trypsin 0.25%: EDTA 0.02% (1:3), split ratio 1:5, optimal population density  $5.0 \times 10^3$ - $1.0 \times 10^4$  cells/cm<sup>2</sup>

cryoconservation - growth medium, 8%DMSO,  $1.0$ - $2.0 \times 10^6$  cells/ml in ampule

**Viability after cryoconservation:** 95% (0 passage, dye trypan blue)

**Sterility:** tests for bacteria, fungi and mycoplasma were negative

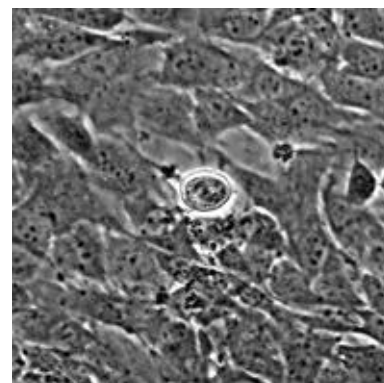
**Species:** karyological and isoenzymological (LDH, G6PD) analysis

**Karyology:**  $2n=40$ , variability in the range between 70-80 chromosomes, modal number of chromosomes 74-76, number of markers - 1-3 metacentric chromosomes (routine dye), number of polyploid cells 5.0%.

**Other properties:** secretion of extracellular matrix protein for adhesion of keratinocytes and growth factors for stimulation of keratinocyte proliferation.

**Applications:** feeder for cultivation of epithelial cells.

**Collections:** SPBIC



## 3T6 Swiss albino

**Origin:** Swiss mouse, embryo

J. Cell Biol. 1963. 17: 299-313; Nature 1966. 212: 631-633.

**Morphology:** fibroblast-like

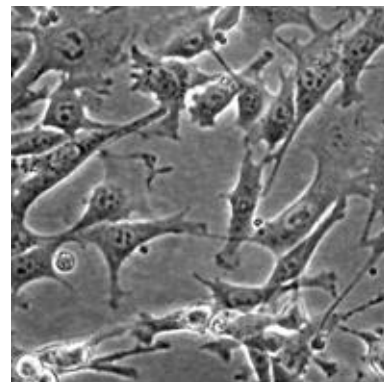
**Mode of cultivation:** monolayer

**Conditions for cultivation:** medium - DMEM

serum - FBS 10%

subculture procedure - cells detach from flask using trypsin 0.25%: EDTA 0.02% (1:1), split ratio 1:5 - 1:8, optimal population density  $5.0 \times 10^3$  -  $1.0 \times 10^4$  cells/cm<sup>2</sup>

cryoconservation - growth medium, 10% DMSO,  $1.0 \times 10^6$  cells/ml in ampule



**Viability after cryoconservation:** 72% (0 passage, dye trypan blue)

**Sterility:** tests for bacteria, fungi and mycoplasma were negative

**Species:** karyological and isoenzymological (LDH, G6PD) analysis

**Karyology:**  $2n=40$ , variability in the range between 64-84 chromosomes, modal number of chromosomes 70-72, some cells have large submetacentric chromosome and microchromosomes (routine dye).

**Plating efficiency:** 32 %.

**Other properties:** virus susceptibility: herpes simplex, vaccinia, vesicular stomatitis (Indiana), pseudorabies.

Collagen and hyaluronic acid secretion.

**Applications:** differentiation, proliferation study.

**Collections:** ATCC CCL 96; ECACC 86120801; SPBIC.

**Origin:** mouse CC57W, rhabdomyosarcoma induced in vivo by methylcholanthrene  
Submitted in Institute of Cytology RAS 1977.

**Morphology:** fibroblast-like

**Mode of cultivation:** monolayer

**Conditions for cultivation:** medium - DMEM

serum - FBS 10%

subculture procedure - cells detach  
from flask using trypsin 0.25%: EDTA  
0.02% (1:3), split ratio 1:5.

cryoconservation - growth medium,  
10% DMSO,  $1.0 \times 10^6$  cells/ml in ampule

**Viability after cryoconservation:** 91% (0 passage,  
dye trypan blue)

**Sterility:** tests for bacteria, fungi and mycoplasma were negative

**Species:** karyological, isoenzymological (LDH) and immunofluorescent analysis.

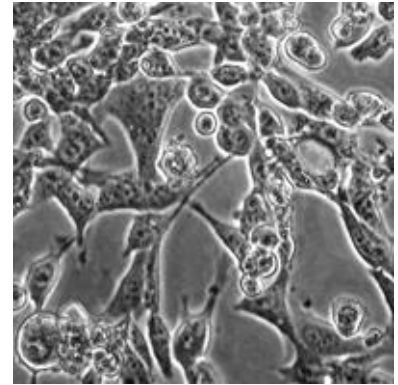
**Karyology:**  $2n = 40$ , variability in the range between 46-63 chromosomes, modal  
number of chromosomes 54-56, number of markers - 1, large metacentric  
chromosomes (routine dye), 1-3 microchromosomes in the most cells, number of  
polyploid cells 28%.

**Plating efficiency:** 88 %

**Tumorigenicity:** tumorigenic in syngeneic mouse.

**Applications:** tumorigenicity, cell biology.

**Collections:** SPBIC.



**Origin:** mouse C3H/An, subcutaneous adipose connective tissue, derived from NCTC 929.

Proc.Natl.Acad.Sci. 1963. 50: 568; Nature 1964. 202: 1142; Am.J.Human Gen. 1974. 26: 273.

**Morphology:** fibroblast-like

**Mode of cultivation:** monolayer

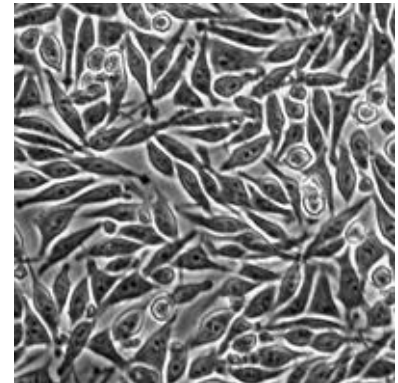
**Conditions for cultivation:** medium - DMEM

serum - FBS 10%

subculture procedure - cells detach from flask using trypsin 0.25%: EDTA 0.02%

(1:1), split ratio 1:3 - 1:10, optimal population density  $1.0-5.0 \times 10^4$  cells/cm<sup>2</sup>

cryoconservation - growth medium, 10% DMSO,  $1.0 \times 10^6$  cells/ml in ampule



**Viability after cryoconservation:** 95% (0 passage, dye trypan blue)

**Sterility:** tests for bacteria, fungi and mycoplasma were negative

**Species:** karyological and isoenzymological (LDH, G6PD) analysis

**Karyology:**  $2n=40$ , variability in the range between 52-57 chromosomes, modal number of chromosomes 54-55, number of markers - 1 (routine dye), number of polyploid cells 1.8%.

**Other properties:** deficient in hypoxanthine phosphoribosyltransferase, resistant to 8-azaguanine and 6-thioguanine

May be heterozygous for the ability to synthesize active inosinic acid phosphorylase.

**Applications:** metabolism, genetics of somatic cells.

**Collections:** ATCC CRL 6319; ECACC 84011426; SPBIC.



**Origin:** Chinese hamster, lung, clone of subline A-23 of cell line DON.

Bioch.Genet. 1972. 7: 33; DAN Russ. 1982. 267. 6: 1496-1498; Tsytologya, 1985. 27. 4: 467-475.

**Morphology:** fibroblast-like

**Mode of cultivation:** monolayer

**Conditions for cultivation:** medium - F10

serum - FBS 10%

subculture procedure - cells detach from flask using trypsin 0.25%: EDTA 0.02% (1:3), split ratio 1:2 - 1:3

cryoconservation - growth medium, 10% DMSO,  $1.0 \times 10^6$  cells/ml in ampule

**Viability after cryoconservation:** 98% (0 passage, dye trypan blue)

**Sterility:** tests for bacteria, fungi and mycoplasma were negative

**Species:** karyological and isoenzymological (LDH, G6PD) analysis

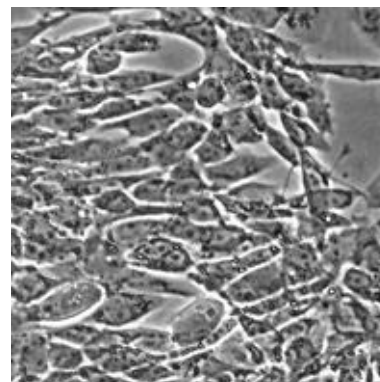
**Karyology:**  $2n=22$ , variability in the range between 30-48 chromosomes, modal number of chromosomes 41-44, number of markers - 8 - in the most cells (differential dye).

**Other properties:**

deficient in thymidine kinase, resistant to BUdR.

**Applications:** cell biology, genetics of somatic cells.

**Collections:** SPBIC.



**Origin:** Chinese hamster, peritoneal cells, fibrosarcoma, derived from B14FAF28-G3. Science 1961. 133: 1600; Tex.Rep.Biol. a Med. 1965. 23: 231.

**Morphology:** fibroblast-like

**Mode of cultivation:** monolayer

**Conditions for cultivation:** medium - DMEM

serum - FBS 10%

subculture procedure - cells detach from flask using trypsin 0.25%: EDTA 0.02% (1:3), split ratio 1:4 – 1:8

cryoconservation - growth medium, 5% DMSO,  $1.0 \times 10^6$  cells/ml in ampule

**Viability after cryoconservation:** 94% (0 passage, dye trypan blue)

**Sterility:** tests for bacteria, fungi and mycoplasma were negative

**Species:** karyological, isoenzymological (LDH, G6PD) and immunofluorescent analysis

**Karyology:**  $2n = 22$ , variability in the range between 19-25 chromosomes, modal number of chromosomes 22, pseudodiploid, a dicentric chromosome was observed in some cells.

**Plating efficiency:** 46 %.

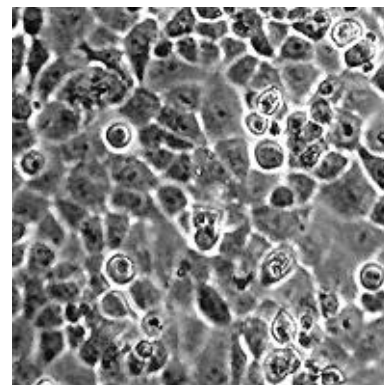
**Other properties:**

virus susceptibility: vesicular stomatitis (Indiana).

deficient in thymidine kinase, resistant to 5 - iododeoxyuridine

**Applications:** genetics, cell biology.

**Collections:** ATCC CCL 14.1; SPBIC.



## BALB/3T3 clone A31

**Origin:** mouse BALB/c, embryo.

J.Cell Physiol. 1968. 72: 141-148; Virology 1969. 38: 174-202; Science 1968. 162: 1024-1026; Exp.Cell Res. 1970. 59: 137.

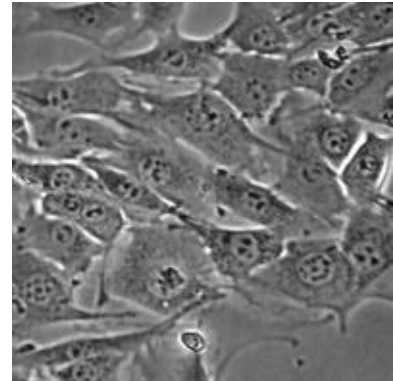
**Morphology:** fibroblast-like

**Mode of cultivation:** monolayer

**Conditions for cultivation:** medium - DMEM  
serum - FBS 10%

subculture procedure - cells detach from flask using trypsin 0.25%: EDTA 0.02% (1:3), split ratio 1:3 - 1:6, optimal population density  $3.0 \times 10^3$  -  $2.0 \times 10^4$  cells/cm<sup>2</sup>

cryoconservation - growth medium, 7.5% DMSO,  $1.0 \times 10^6$  cells/ml in ampule



**Viability after cryoconservation:** 90% (0 passage, dye trypan blue)

**Sterility:** tests for bacteria, fungi and mycoplasma were negative

**Species:** karyological and isoenzymological (LDH, G6PD) analysis

**Karyology:**  $2n = 40$ , variability in the range between 55-84 chromosomes, modal number of chromosomes 68-74, number of polyploid cells 3.0%.

**Plating efficiency:** 20 %.

**Other properties:** virus susceptibility: herpes simplex, vesicular stomatitis, coronavirus, SV 40, vaccinia, polyoma.

Contact inhibition of growth (by density  $2.0$ - $2.5 \times 10^5$  cells/cm<sup>2</sup>).

**Applications:** virology, replication, tumorigenicity.

**Collections:** ATCC CCL 163; ECACC 86110401; MWIIW; SPBIC.

**Origin:** mouse C3H, smooth muscle-like cells from brain tumor induced in vivo by ethyl nitrosoethylurea.

J.Cell Biol. 1974. 61: 318-413; J.Biol.Chem. 1977. 252: 2143-2153.

**Morphology:** fibroblast-like

**Mode of cultivation:** monolayer

**Conditions for cultivation:** medium - DMEM

serum - FBS 20%

subculture procedure - cells detach from flask using trypsin 0.25%: EDTA 0.02% (1:3), split ratio 1:3 - 1:6, optimal population density  $1.0-5.0 \times 10^4$  cells/cm<sup>2</sup>

cryoconservation - growth medium, 8% DMSO,  $1.0 \times 10^6$  cells/ml in ampule

**Viability after cryoconservation:** 95% (0 passage, dye trypan blue)

**Sterility:** tests for bacteria, fungi and mycoplasma were negative

**Species:** karyological analysis

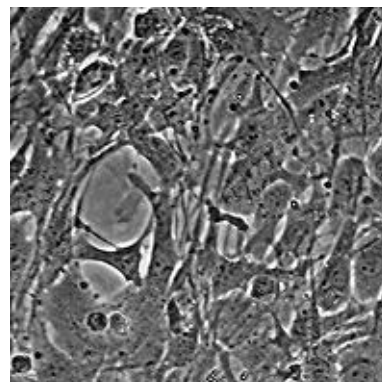
**Karyology:**  $2n=40$ , variability in the range between 60-76 chromosomes, modal number of chromosomes 64-67, number of polyploid cells 8%.

**Other properties:** synthesis of adenylate and creatine phosphokinases, acetylcholine receptors.

Possess many properties characteristic of smooth muscle.

**Applications:** acetylcholine receptors study.

**Collections:** ATCC CRL 1443; ECACC 86093001; SPBIC.



**Origin:** African green monkey, kidney.

Arch.Gesamte Virusforsch. 1970. 32: 389; Health Lab. Sci. 1974. 110: 275; Append. Environ.Microbiol. 1986. 51: 790.

**Morphology:** epithelial-like

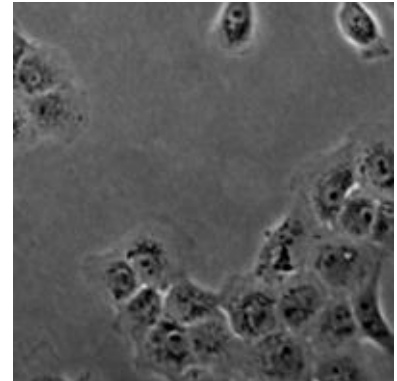
**Mode of cultivation:** monolayer

**Conditions for cultivation:** medium - DMEM

serum - FBS 10%

subculture procedure - cells detach from flask using trypsin 0.25%: EDTA 0.02% (1:3), split ratio 1:3 - 1:6, optimal population density  $5.0 \times 10^3$  -  $2.0 \times 10^4$  cells/cm<sup>2</sup>

cryoconservation - growth medium, 10% DMSO,  $1.0 \times 10^6$  cells/ml in ampule



**Viability after cryoconservation:** 95% (0 passage, dye trypan blue)

**Sterility:** tests for bacteria, fungi and mycoplasma were negative

**Species:** karyological and isoenzymological (LDH and G6PD) analysis

**Karyology:**  $2n=60$ , variability in the range between 58-68 chromosomes, modal number of chromosomes 61-62, number of markers - 1-2, small submetacentric chromosome with secondary constriction (routine dye)

**Other properties:** virus susceptibility: poliovirus 1, 2, 3; ECHO 3, 6, 7, 9, 11, 12, 27; Coxsackie A9, B1, B2, B3; reovirus; rotavirus SA 11.

**Applications:** virology, chlamidia growth substrate.

**Collections:** ECACC 90092601; MWIIW; SPBIC.

**Origin:** Syrian hamster, kidney

Virology 1962. 16: 147-151; J.Natl.Cancer Inst. 1963. 30: 795-811.

**Morphology:** fibroblast-like

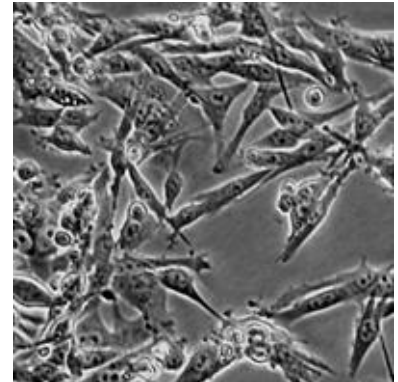
**Mode of cultivation:** monolayer

**Conditions for cultivation:** medium - DMEM

serum - FBS 10%

subculture procedure - cells detach from flask using trypsin 0.25%: EDTA 0.02% (1:3), split ratio 1:3 - 1:6, optimal population density  $2.0-4.0 \times 10^4$  cells/cm<sup>2</sup>

cryoconservation - growth medium, 5 – 10% DMSO,  $1.0-1.5 \times 10^6$  cells/ml in ampule



**Viability after cryoconservation:** 92% (0 passage, dye trypan blue)

**Sterility:** tests for bacteria, fungi and mycoplasma were negative

**Species:** karyological and isoenzymological (LDH and G6PD) analysis.

**Karyology:**  $2n=44$ , variability in the range between 44-52 chromosomes, modal number of chromosomes 49-50, number of markers - 1 large metacentric chromosome (routine dye), 7 markers (differential dye), number of polyploid cells 5.1%

**Other properties:** virus susceptibility: pseudorabies, vaccinia, herpes simplex, reovirus 3; vesicular stomatitis (Indiana), rubella, adenovirus 25, foot-and-mouth disease virus, Coxsackie, rabies, arboviruses..

**Applications:** virology, transformation, cell biology.

**Collections:** ATCC CCL 10; ECACC 85011433; SPBII; SPBIC, MWIEV.

**Origin:** mouse, hepatoma

J.Cell Sci. 1979. 35: 267; Exp.Cell Res. 1980. 125: 305.

**Morphology:** fibroblast-like

**Mode of cultivation:** monolayer

**Conditions for cultivation:** medium - DMEM

serum - FBS 10%

subculture procedure - cells detach from flask using trypsin 0.25%: EDTA 0.02% (1:3), split ratio 1:3 - 1:5.

cryoconservation - growth medium, 10% DMSO,  $1.0 \times 10^6$  cells/ml in ampule

**Viability after cryoconservation:** 86% (0 passage, dye trypan blue)

**Sterility:** tests for bacteria, fungi and mycoplasma were negative

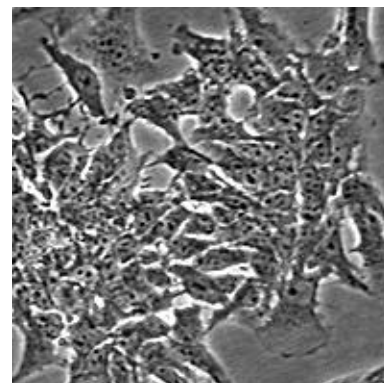
**Species:** karyological and isoenzymological (LDH and G6PD) analysis.

**Karyology:**  $2n = 40$ , variability in the range between 62-68 chromosomes, modal number of chromosomes 65-66, number of markers - 1-3, large meta- and submetacentric chromosomes, the most cells have small metacentric chromosomes (routine dye), number of polyploid cells 2.4%.

**Other properties:** deficient in hypoxanthine phosphoribosyltransferase, resistant to 8 - azaguanine and 6 - thioguanine

**Applications:** somatic cell genetics

**Collections:** SPBIC



**Origin:** rat, glioma induced in vivo by N-nitrosomethylurea, monoclonal cell line. Science 1968. 161: 370; Fed.Proc. 1968. 27: 720; Atlas of chromosomes of human and animal cell lines, S.E. Mamaeva, 2002. Moscow, Scientific world

**Morphology:** fibroblast-like

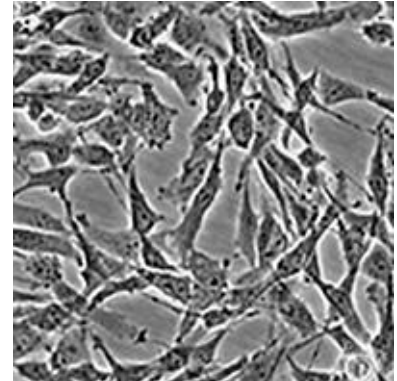
**Mode of cultivation:** monolayer

**Conditions for cultivation:** medium - F10

serum - HS 15%/FBS 2.5%

subculture procedure - cells detach from flask using trypsin 0.25%: EDTA 0.02% (1:3), split ratio 1:3 - 1:6, optimal population density  $1.0-3.0 \times 10^5$  cells/cm<sup>2</sup>

cryoconservation - growth medium, 7.5%DMSO,  $1.0-2.0 \times 10^6$  cells/ml in ampule



**Viability after cryoconservation:** 93% (0 passage, dye trypan blue)

**Sterility:** tests for bacteria, fungi and mycoplasma were negative

**Species:** karyological, immunofluorescent and isoenzymological (LDH, G6PD) analysis

**Karyology:**  $2n=42$ , variability in the range between 39-44 chromosomes, modal number of chromosomes 42, normal rat karyotype (42, XY), cells, containing 43 chromosomes have 1 marker (differential dye).

**Plating efficiency:** 26%.

**Tumorigenicity:** tumorigenic in albino rat

**Other properties:** virus susceptibility: pseudorabies, vesicular stomatitis (Indiana), herpes simplex, vaccinia.

S 100 protein production

**Applications:** biochemistry, virology, differentiation, tumorigenicity.

**Collections:** ATCC CCL 107; ECACC 85040101; ICLC ATL 95007; SPBIC.



**Origin:** mouse C3H, leg muscle.

Nature 1977. 270: 725-727; Science 1985. 230: 758-766.

**Morphology:** myoblast-like

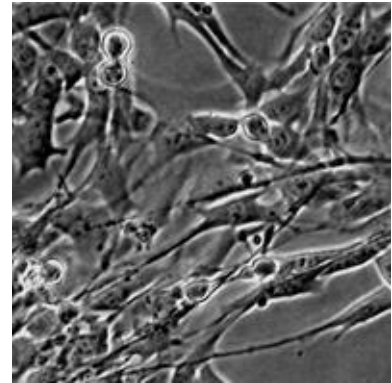
**Mode of cultivation:** monolayer

**Conditions for cultivation:** medium - DMEM

serum - FBS 10%

subculture procedure - cells detach from flask using trypsin 0.25%: EDTA 0.02% (1:3), split ratio 1:3 - 1:8, optimal population density  $2.0-4.0 \times 10^4$  cells/cm<sup>2</sup>

cryoconservation - growth medium, 10% DMSO,  $1.0 \times 10^6$  cells/ml in ampule



**Viability after cryoconservation:** 90-95% (0 passage, dye trypan blue)

**Sterility:** tests for bacteria, fungi and mycoplasma were negative.

**Species:** karyological and isoenzymological (LDH, G6PD) analysis

**Karyology:**  $2n=40$ , variability in the range between 73-80 chromosomes, modal number of chromosomes 77-80, number of polyploid cells 0.8%.

**Other properties:** muscle protein expression.

Differentiates producing myotubes.

**Applications:** myogenesis, cell differentiation, cell biology.

**Collections:** ATCC CRL 1772; ECACC 91031101; SPBIC.

## C3H10T1/2 clone 8

**Origin:** mouse C3H, embryo.

Cancer Res. 1973. 33: 3231-3238 и 3239-3249; Nature 1975. 253: 548-549; Virology 1975. 65: 392-409.

**Morphology:** fibroblast-like

**Mode of cultivation:** monolayer

**Conditions for cultivation:** medium - DMEM  
serum - FBS 10%

subculture procedure - cells detach from flask using trypsin 0.25%: EDTA 0.02% (1:3), split ratio 1:3 - 1:6, optimal population density  $2.0 \times 10^3$  cells/cm<sup>2</sup>

cryoconservation - growth medium, 10% DMSO,  $1.5 \times 10^6$  cells/ml in ampule

**Viability after cryoconservation:** 92% (0 passage, dye trypan blue)

**Sterility:** tests for bacteria, fungi and mycoplasma were negative

**Species:** isoenzymological (LDH, G6PD) analysis

**Karyology:**  $2n=40$ , variability in the range between 66-82 chromosomes, modal number of chromosomes 80, number of markers - 16 (differential dye).

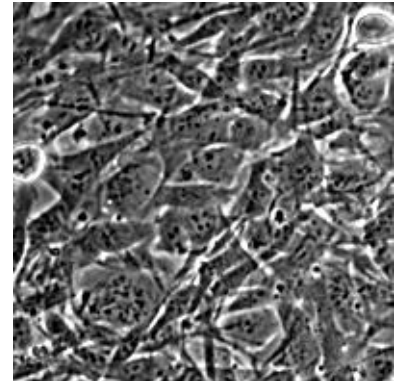
**Plating efficiency:** 30%.

**Tumorigenicity:** non tumorigenic

**Other properties:** contact inhibition of growth

**Applications:** tumorigenicity, transformation, transfection, cell biology.

**Collections:** ATCC CCL 226; ECACC 86060303; SPBIC.



**Origin:** Chinese hamster, ovary, clone CHO.

J.Exp.Med. 1958. 108: 945; Proc. Natl.Acad.Sci. USA 1968. 60: 1275.

**Morphology:** epithelial-like

**Mode of cultivation:** monolayer

**Conditions for cultivation:** medium - F12

serum - FBS 10%

subculture procedure - cells detach from flask using trypsin 0.25%: EDTA 0.02% (1:3), split ratio 1:3 - 1:8, optimal

population density  $1.0-2.0 \times 10^4$  cells/cm<sup>2</sup>

cryoconservation - growth medium, 10% DMSO,  $1.0-2.0 \times 10^6$  cells/ml in ampule

**Viability after cryoconservation:** 99% (0 passage, dye trypan blue)

**Sterility:** tests for bacteria, fungi and mycoplasma were negative

**Species:** karyological, isoenzymological (LDH, G6PD) and immunofluorescent analysis

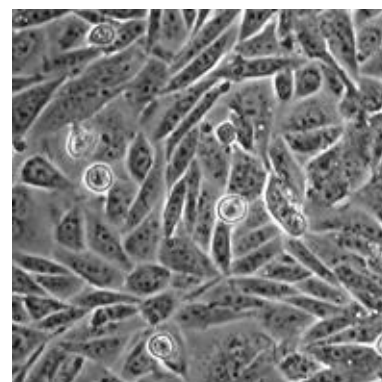
**Karyology:**  $2n=22$ , variability in the range between 16-22 chromosomes, modal number of chromosomes 20, number of markers - 11 (differential dye), number of polyploid cells 7.4%

**Plating efficiency:** 90%.

**Other properties:** virus susceptibility: vesicular stomatitis (Indiana), Getah arbovirus. Absence of the gene for proline synthesis, requirement of proline for growth.

**Applications:** somatic cells genetics, cell biology, virology.

**Collections:** ATCC CCL 61; ECACC 85051005; DSM ACC 110; ESCC; SPBIC.



## Clone M-3

**Origin:** mouse F<sub>1</sub> (CxDBA), clone from melanoma Cloudman S91.  
Science 1966. 154: 1186.

**Morphology:** epithelial-like

**Mode of cultivation:** monolayer

**Conditions for cultivation:** medium - F12

serum - FBS 10%

subculture procedure - cells detach from flask using trypsin 0.25%: EDTA 0.02% (1:3), split ratio 1:2 - 1:3, optimal

population density  $2.0-3.0 \times 10^4$  cells/cm<sup>2</sup>

cryoconservation - growth medium, 10% DMSO,  $2.0-3.0 \times 10^6$  cells/ml in ampule

**Viability after cryoconservation:** 94% (0 passage, dye trypan blue)

**Sterility:** tests for bacteria, fungi and mycoplasma were negative

**Species:** karyological and isoenzymological (LDH, G6PD) analysis

**Karyology:**  $2n=40$ , variability in the range between 76-86 chromosomes, modal number of chromosomes 83, number of markers - 2 (routine dye), some cells have microchromosomes.

**Plating efficiency:** less than 1%.

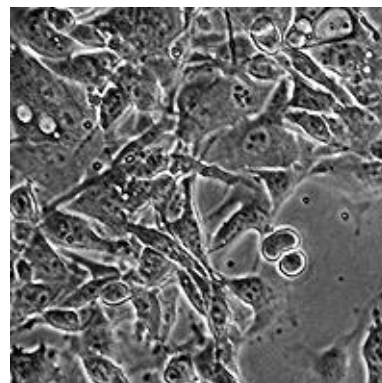
**Tumorigenicity:** tumorigenic in syngeneic animals

virus susceptibility: herpes simplex, vaccinia, pseudorabies, vesicular stomatitis (Indiana).

Melanin production for at least 33 passages

**Applications:** virology, tumorigenicity, cell biology.

**Collections:** ATCC CCL 53.1; ECACC 87081806; SPBIC.



**Origin:** African green monkey, kidney.

Proc.Natl.Acad.Sci. 1964. 53: 53; Virology 1965. 27: 453.

**Morphology:** fibroblast-like

**Mode of cultivation:** monolayer

**Conditions for cultivation:** medium - DMEM

serum - FBS 10%

subculture procedure - cells detach from flask using trypsin 0.25%: EDTA 0.02%

(1:3), split ratio 1:3 - 1:6, optimal

population density  $2.0-4.0 \times 10^4$  cells/cm<sup>2</sup>

cryoconservation - growth medium,

5%DMSO,  $1.0 \times 10^6$  cells/ml in ampule

**Viability after cryoconservation:** 85% (0 passage, dye trypan blue)

**Sterility:** tests for bacteria, fungi and mycoplasma were negative

**Species:** karyological and isoenzymological (LDH, G6PD) analysis

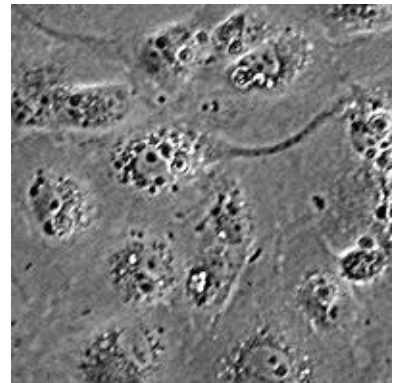
**Karyology:**  $2n=60$ , variability in the range between 56-61 chromosomes, modal number of chromosomes 60, number of markers - 4-5 (differential dye), number of polyploid cells 4.4%.

**Plating efficiency:** 27%.

**Other properties:** virus susceptibility: poliovirus 1, herpes simplex, Eastern equine encephalitis, Western equine encephalitis, California encephalitis, SV 40.

**Applications:** virology.

**Collections:** ATCC CCL 70; ECACC 87032605; SPBII; MWIIV; SPBIC.



**Origin:** rat, embryo, fibroblasts transformed by adenovirus 5.  
Mol.Biol. (Russ.) 1979. 13: 292.

**Morphology:** fibroblast-like

**Mode of cultivation:** monolayer

**Conditions for cultivation:** medium - EMEM

serum - FBS 10%

subculture procedure - cells detach  
from flask using EDTA 0.02%, split ratio  
1:3

cryoconservation - growth medium, 10%  
DMSO,  $2.0 \times 10^6$  cells/ml in ampule

**Viability after cryoconservation:** 85% (0 passage, dye  
trypan blue)

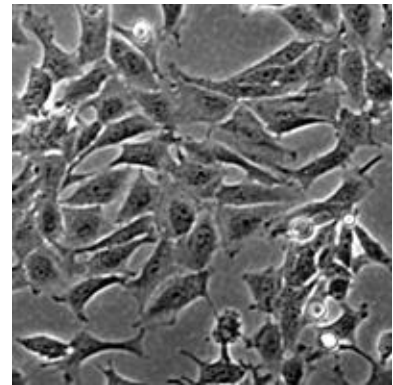
**Sterility:** tests for bacteria, fungi and mycoplasma were negative

**Species:** karyological and isoenzymological (LDH, G6PD) analysis

**Karyology:**  $2n=42$ , variability in the range between 55-67 chromosomes, modal  
number of chromosomes 64-65, number of polyploid cells 3.0%.

**Applications:** cell biology.

**Collections:** SPBIC



**Origin:** Chinese hamster, ovary, clone of CHO.

Submitted from Columbia University, New York, USA, 1984; Digest "Cell Cultures" 2015. 31: 46 – 54.

**Morphology:** epithelial-like

**Mode of cultivation:** monolayer

**Conditions for cultivation:** medium - F12

serum - FBS 10%

subculture procedure - cells detach from flask using trypsin 0.25%: EDTA 0.02% (1:3), split ratio 1:3

cryoconservation - growth medium, 10% DMSO,  $1.0-2.0 \times 10^6$  cells/ml in ampule

**Viability after cryoconservation:** 90% (0 passage, dye trypan blue)

**Sterility:** tests for bacteria, fungi and mycoplasma were negative

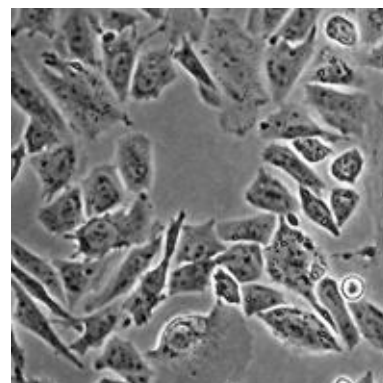
**Species:** karyological analysis

**Karyology:**  $2n=22$ , variability in the range between 18-22 chromosomes, modal number of chromosomes 20, number of markers - 2 large metacentric chromosomes (routine dye), number of markers – 14 (defferential dye), number of polyploid cells 1.2%.

**Other properties:** dihydrofolate reductase deficient, requires hypoxanthine or adenine, glycine, thymidine and proline.

**Applications:** biochemistry, cell biology.

**Collections:** SPBIC



**Origin:** mouse C57BL/6N, lymphoma induced by dimethyl-benzanthracene (ascitic fluid).

Br.J.Cancer 1950. 4:372; Cancer Res. 1965. 25: 813; J.Natl.Cancer Inst. 1972. 48: 265; J.Jmmunol. 1972. 108:1146; J.Jmmunol.1973. 110: 1470.

**Morphology:** lymphoid

**Mode of cultivation:** suspension

**Conditions for cultivation:** medium - RPMI 1640

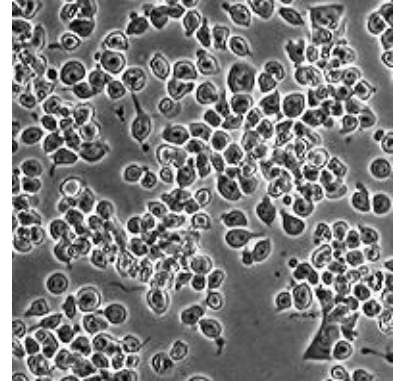
serum - FBS 10%

subculture procedure - optimal

population density  $3.0-9.0 \times 10^5$  cells/ml

cryoconservation - growth medium,

10% DMSO,  $5.0-6.0 \times 10^6$  cells/ml in ampule



**Viability after cryoconservation:** 70% (0 passage, dye trypan blue)

**Sterility:** tests for bacteria, fungi and mycoplasma were negative

**Species:** karyological and isoenzymological (LDH, G6PD) analysis

**Karyology:**  $2n=40$ , variability in the range between 36-40 chromosomes, modal number of chromosomes 38 and 40, number of markers - 3-4 (routine dye), number of polyploid cells 2.2%.

**Other properties:** antigens expressed by these cells include: G, a surface antigen induced by leukemia type G virus; H-2<sup>b</sup> and Thy-1,2.

These cells do not bear TL antigen or surface immunoglobulin.

Resistant to cortisol and dexamethasone.

Sensitive to PHA.

**Applications:** virology, tumorigenicity, biotechnology (IL-2 and interferon production).

**Collections:** ATCC TIB 39; ECACC 85022105; SPBIC.



**Origin:** mouse C57Bl, glioblastoma induced by dimethylbenzanthracene and then passed in outbred mice.

Tsitologiya, 1977. 19. 1: 95-100.

**Morphology:** fibroblast-like

**Mode of cultivation:** monolayer

**Conditions for cultivation:** medium - EMEM

serum - FBS 10%

subculture procedure - cells detach from flask using trypsin 0.25%: EDTA 0.02% (1:3), split ratio 1:3 -1:5

cryoconservation - growth medium, 10% DMSO,  $1.0 \times 10^6$  cells/ml in ampule

**Viability after cryoconservation:** 94% (0 passage, dye trypan blue)

**Sterility:** tests for bacteria, fungi and mycoplasma were negative

**Species:** karyological, isoenzymological (LDH, G6PD) and immunofluorescent analysis

**Karyology:**  $2n=40$ , variability in the range between 57-64 chromosomes, modal number of chromosomes 59-60, number of markers-3-5 large-sized metacentric and 1 middle acrocentric with secondary constriction (routine dye), number of polyploid cells 1.5%.

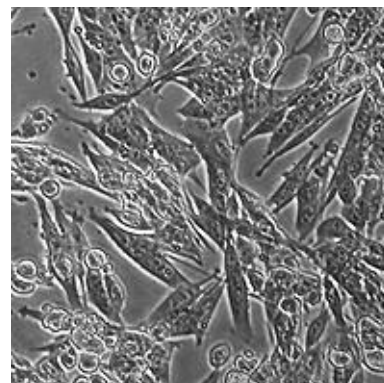
**Plating efficiency:** 60%

**Tumorigenicity:** tumorigenic in outbred mice

**Other properties:** muscarinic and nicotinic receptors for acetylcholine and receptors for diazepam.

**Applications:** neurooncology, cell biology.

**Collections:** SPBIC.



**Origin:** mouse line 129, testicular teratocarcinoma  
 Proc. Natl. Acad. Sci. USA 1973. 70: 3899 – 3903; Cell 1978.15: 393 – 403; Cell 1980. 21: 347 – 355; Atlas of chromosomes of human and animal cell lines, S.E. Mamaeva, 2002. Moscow, Scientific world.

**Morphology:** epithelial-like

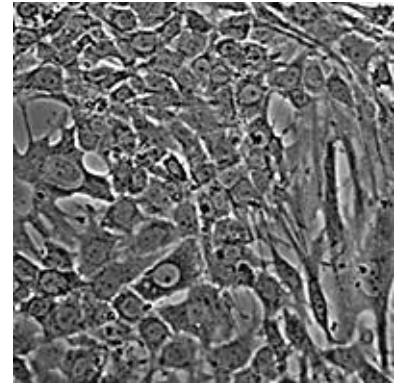
**Mode of cultivation:** monolayer

**Conditions for cultivation:** medium - DMEM

serum - FBS 10%

subculture procedure - culture surface are coated with 0.1% gelatin, cells detach from flask using trypsin 0.25%: EDTA 0.02% (1:3), split ratio 1:3 - 1:6, optimal population density  $2.0-4.0 \times 10^4$  cells/cm<sup>2</sup>

cryoconservation - growth medium, 5% DMSO,  $1.5 \times 10^6$  cells/ml in ampule



**Viability after cryoconservation:** 85 % (0 passage, dye trypan blue)

**Sterility:** tests for bacteria, fungi and mycoplasma were negative

**Species:** karyological analysis

**Karyology:**  $2n = 40$ , variability in the range between 37-41 chromosomes, modal number of chromosomes 39, number of markers - 8 (differential dye), number of polyploid cells 0.8%.

**Other properties:** undergo very limited differentiation under normal culture conditions; Induction of differentiation into parietal endoderm in the presence of retinoic acid and dibutyryl cyclic AMP;

Synthesis of plasminogen activator, laminin, type IV collagen, low levels alkaline phosphatase and lactate dehydrogenase.

**Applications:** cell biology, differentiation, tumorigenicity.

**Collections:** ATCC CRL 1720; ECACC 85060401; SPBIC.

**Origin:** bovine, embryo, trachea.

Folia Biol. 1975. 21: 117.

**Morphology:** epithelial-like

**Mode of cultivation:** monolayer

**Conditions for cultivation:** medium - DMEM

serum - FBS 10%

subculture procedure - cells detach from flask using trypsin 0.25%: EDTA 0.02% (1:3), split ratio 1:3 -1:5.

cryoconservation - growth medium, 8-10% DMSO,  $1.0 \times 10^6$  cells/ml in ampule

**Viability after cryoconservation:** 98% (0 passage, dye trypan blue)

**Sterility:** tests for bacteria, fungi and mycoplasma were negative

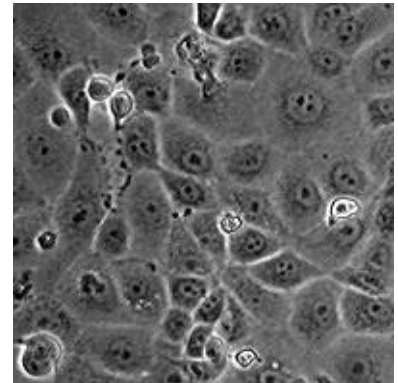
**Species:** karyological and isoenzymological (LDH, G6PD) analysis

**Karyology:**  $2n = 60$ , variability in the range between 42-53 chromosomes, modal number of chromosomes 48-49, number of polyploid cells 0.2%.

**Other properties:** virus susceptibility: vesicular stomatitis, IBR, parainfluenza 3.

**Applications:** virology.

**Collections:** MWIIV; SPBIC.



**Origin:** rat, pituitary tumor.

Endocrinology 1968. 82: 342; J.Cell Biol. 1969. 43: 432; In Vitro 1970. 60: 180.

**Morphology:** epithelial-like

**Mode of cultivation:** monolayer

**Conditions for cultivation:** medium - F10

serum - HS 15%, FBS 2.5%

subculture procedure - cells detach

from flask using trypsin 0.25%: EDTA

0.02% (1:3), split ratio 1:3 - 1:4, optimal

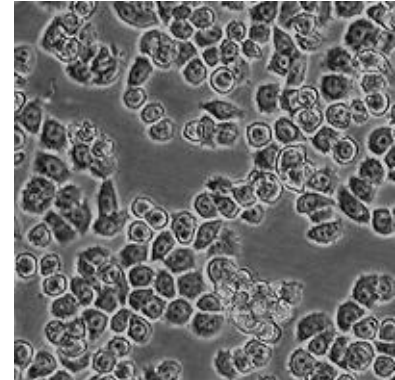
population density  $2.0-4.0 \times 10^4$

cells/cm<sup>2</sup>

cryoconservation - growth medium,

10% DMSO,  $1.0 - 2.0 \times 10^6$  cells/ml in

ampule



**Viability after cryoconservation:** 85% (0 passage, dye trypan blue)

**Sterility:** tests for bacteria, fungi and mycoplasma were negative

**Species:** karyological and isoenzymological (LDH, G6PD) analysis

**Karyology:**  $2n=42$ , variability in the range between 40-75 chromosomes without modal number, number of markers - 2 dicentrics (routine dye), number of polyploid cells 0.6%.

**Plating efficiency:** less than 1%.

**Tumorigenicity:** tumorigenic in syngeneic animals

**Other properties:** virus susceptibility: vesicular stomatitis (Indiana), herpes simplex.

Growth hormone, prolactin, somatotrophin secretion.

**Applications:** endocrinology, cell biology.

**Collections:** ATCC CCL 82.1; ECACC 87012603; ICLC ATL 96003; SPBIC.

**Origin:** Syrian hamster, kidney

S.Afr.J.Med.Sci. 1963. 28: 81.

**Morphology:** epithelial-like

**Mode of cultivation:** monolayer

**Conditions for cultivation:** medium - DMEM

serum - FBS 10%

subculture procedure - cells detach

from flask using trypsin 0.25%: EDTA

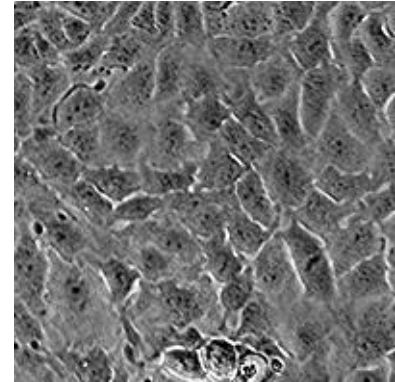
0.02% (1:3), split ratio 1:5 - 1:6, optimal

population density  $2.0-4.0 \times 10^4$

cells/cm<sup>2</sup>

cryoconservation - growth medium,

10% DMSO,  $1.0 \times 10^6$  cells/ml in ampule



**Viability after cryoconservation:** 86% (0 passage, dye trypan blue)

**Sterility:** tests for bacteria, fungi and mycoplasma were negative

**Species:** karyological and isoenzymological (LDH, G6PD) analysis

**Karyology:**  $2n=44$ , variability in the range between 48-58 chromosomes, modal number of chromosomes 52-53 and 56, number of polyploid cells 5.0%.

**Plating efficiency:** 50%.

**Tumorigenicity:** tumorigenic in hamster

**Other properties:** virus susceptibility: vesicular stomatitis, arboviruses, Coxsackie A4, A8, B1, herpes simplex, smallpox, Asian strain influenza, influenza, alpha viruses.

**Applications:** virology.

**Collections:** ATCC CCL 15; ECACC 90102522; MWIHW; SPBIC.

**Origin:** rat Buffalo, hepatoma induced by N,N'-2,7-fluorenylenebis-2,2,2-trifluoroacetamide, ascitic fluid.

Proc.Natl.Acad.Sci. 1966. 56: 296; ATLA 1988. 16: 32.

**Morphology:** epithelial-like

**Mode of cultivation:** monolayer

**Conditions for cultivation:** medium - DMEM

serum - FBS 10%

subculture procedure - cells detach from flask using trypsin 0.25%: EDTA 0.02% (1:3), split ratio 1:5

cryoconservation - growth medium, 10% DMSO,  $1.0 \times 10^6$  cells/ml in ampule

**Viability after cryoconservation:** 90% (0 passage, dye trypan blue)

**Sterility:** tests for bacteria, fungi and mycoplasma were negative

**Species:** karyological and isoenzymological (LDH, G6PD) analysis

**Karyology:**  $2n=42$ , variability in the range between 63-68 chromosomes and 36% of cells have more of 84 chromosomes, modal number of chromosomes 65-67, number of markers - 22 (differential dye)

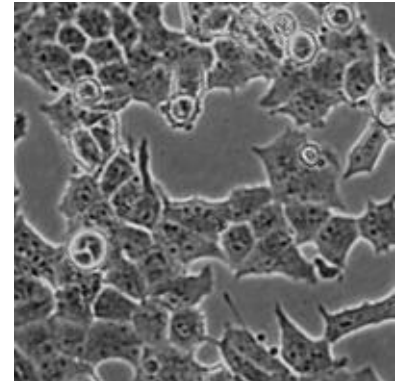
**Plating efficiency:** 60%.

**Tumorigenicity:** tumorigenic in syngeneic animals

**Other properties:** inducible tyrosine aminotransferase.

**Applications:** tumorigenicity, enzymology, cytotoxicity, cell biology.

**Collections:** ICLC ATL 95006; SPBIC.



## Indian Muntjac (M)

**Origin:** muntjac, skin.

Science 1970.168: 1364-1366; Cytogenet.Cell Genet.1979.24: 201-208; Atlas of chromosomes of human and animal cell lines, S.E. Mamaeva, 2002. Moscow, Scientific world.

**Morphology:** fibroblast-like

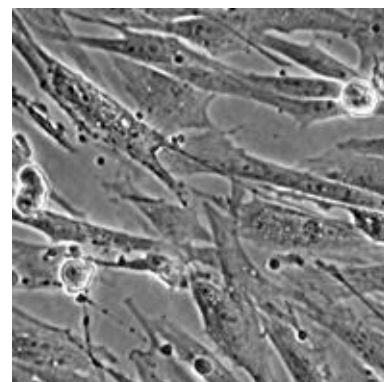
**Mode of cultivation:** monolayer

**Conditions for cultivation:** medium - F10

serum - FBS 20%

subculture procedure - cells detach from flask using trypsin 0.25%: EDTA 0.02% (1:3), split ratio 1:2.

cryoconservation - growth medium, 8-10% DMSO,  $1.0-1.5 \times 10^6$  cells/ml in ampule



**Viability after cryoconservation:** 95% (0 passage, dye trypan blue)

**Sterility:** tests for bacteria, fungi and mycoplasma were negative

**Species:** karyological and isoenzymological (LDH and G6PD) analysis

**Karyology:**  $2n=7$ , variability in the range between 5-12 chromosomes, modal number of chromosomes 7, normal Muntjac karyotype (7, X, Y<sub>1</sub>, Y<sub>2</sub>), number of polyploid cells 3%.

**Plating efficiency:** 29%.

**Other properties:** virus susceptibility: vesicular stomatitis, herpes simplex, vaccinia.

**Applications:** genetics, morphology, virology, cell biology.

**Collections:** ATCC CCL 157; MWIIW; SPBIC.

## Indian Muntjac (MT)

**Origin:** muntjac, skin, subline, spontaneous derived from line M.  
Tsitolgiya. 1988. 31: 807 – 817.

**Morphology:** fibroblast-like

**Mode of cultivation:** monolayer

**Conditions for cultivation:** medium - F10

serum - FBS 20%

subculture procedure - cells detach from flask using trypsin 0.25%: EDTA 0.02% (1:3), split ratio 1:2.

cryoconservation - growth medium, 8-10% DMSO,  $1.0-1.5 \times 10^6$  cells/ml in ampule

**Viability after cryoconservation:** 95% (0 passage, dye trypan blue)

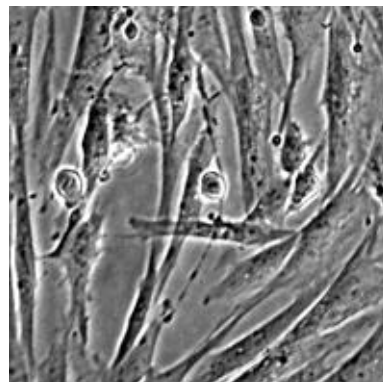
**Sterility:** tests for bacteria, fungi and mycoplasma were negative

**Species:** karyological and isoenzymological (LDH and G6PD) analysis

**Karyology:**  $2n=7$ , variability in the range between 5-12 chromosomes, modal number of chromosomes 9, markers are absent, The difference from normal Muntjac karyotype (7, X, Y<sub>1</sub>, Y<sub>2</sub>) consist of number of homologous chromosomes, number of polyploid cells 3%.

**Applications:** cytogenetics, morphology, cell biology.

**Collections:** SPBIC.





**Origin:** mouse BALB/c, histiocytic sarcoma.

J.Biol.Chem. 1987. 262: 8884; J.Cell Biol. 1988. 106: 657; Proc.Natl.Acad.Sci. 1984. 81: 5430.

**Morphology:** star- and round-shaped

**Mode of cultivation:** monolayer

**Conditions for cultivation:** medium - DMEM

serum - FBS 10%

subculture procedure - cells detach from flask using EDTA 0.02%, split ratio 1:2 - 1:4

cryoconservation - growth medium, 10% DMSO,  $1.0 \times 10^6$  cells/ml in ampule

**Viability after cryoconservation:** 80% (0 passage, dye trypan blue)

**Sterility:** tests for bacteria, fungi and mycoplasma were negative

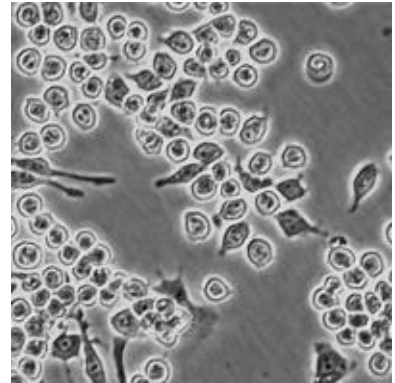
**Species:** karyological analysis

**Tumorigenicity:** tumorigenic in syngeneic animals

**Other properties:** phagocytosis, chemotaxis, antigen presentation.

**Applications:** immunology, cytotoxicity, cell biology.

**Collections:** SPBIC.



**Origin:** rat, sarcoma, derived from cell line Jensen Sarcoma.  
Cancer Res. 1959. 19: 591; Cell 1975. 6: 53-60.

**Morphology:** fibroblast-like

**Mode of cultivation:** monolayer

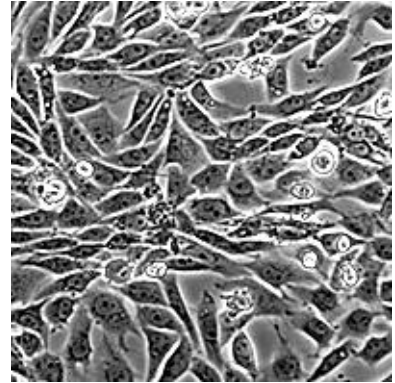
**Conditions for cultivation:** medium - DMEM

serum - FBS 5-10%

other components - NEAA 1%

subculture procedure - cells detach from flask using EDTA 0.02%, split ratio 1:4 - 1:6.

cryoconservation - growth medium, 10% DMSO,  $1.0 \times 10^6$  cells/ml in ampule



**Viability after cryoconservation:** 79% (0 passage, dye trypan blue)

**Sterility:** tests for bacteria, fungi and mycoplasma were negative

**Species:** karyological and isoenzymological (LDH, G6PD) analysis

**Karyology:**  $2n=42$ , variability in the range between 49-61 chromosomes, modal number of chromosomes 52-56, number of markers - 1 middle acrocentric chromosome with gap (routine dye).

**Plating efficiency:** 46%.

**Tumorigenicity:** highly tumorigenic

**Other properties:** requires asparagine for growth

**Applications:** somatic cell genetics, tumorigenicity.

**Collections:** SPBIC

**Origin:** rat, fibroblasts spontaneously transformed in vitro.

Submitted from N.K.Belisheva, Institute of Cytology of the USSR Academy of Sciences, Leningrad, 1976. Dissert. work, 1979. L.

**Morphology:** fibroblast-like

**Mode of cultivation:** monolayer

**Conditions for cultivation:** medium - EMEM

serum - FBS 10%

subculture procedure - cells detach from flask using trypsin 0.25%: EDTA 0.02% (1:3), split ratio 1:4

cryoconservation - growth medium, 5 – 10% DMSO,  $1.0 \times 10^6$  cells/ml in ampule

**Viability after cryoconservation:** 80% (0 passage, dye trypan blue)

**Sterility:** tests for bacteria, fungi and mycoplasma were negative

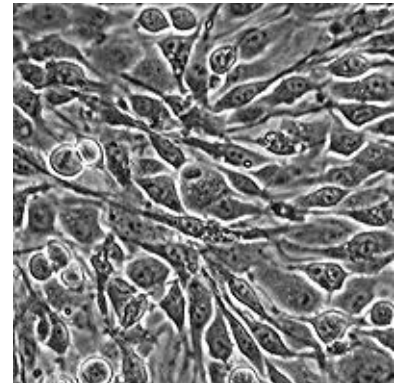
**Species:** karyological and isoenzymological (LDH, G6PD) analysis

**Karyology:**  $2n=42$ , variability in the range between 41-44 chromosomes, modal number of chromosomes 42, 15% of cells have 78-83 chromosomes.

**Tumorigenicity:** highly tumorigenic

**Applications:** cell biology.

**Collections:** SPBIC



**Origin:** rat Wistar, skeletal muscle.

Develop.Biol. 1970. 23: 1-22; Differentiation 1977. 7: 159-166.

**Morphology:** myoblast-like

**Mode of cultivation:** monolayer

**Conditions for cultivation:** medium - DMEM

serum - FBS 10%

subculture procedure - cells detach from flask using trypsin 0.25%: EDTA 0.02% (1:3), split ratio 1:4 - 1:8

cryoconservation - growth medium, 10% DMSO,  $1.0 \times 10^6$  cells/ml in ampule

**Viability after cryoconservation:** 90% (0 passage, dye trypan blue)

**Sterility:** tests for bacteria, fungi and mycoplasma were negative

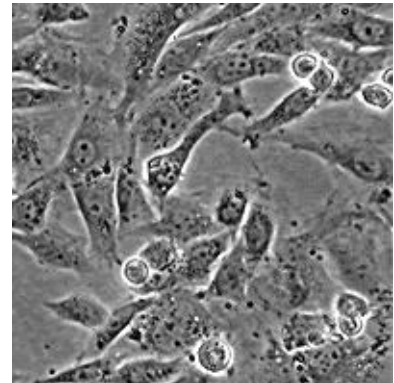
**Species:** karyological and isoenzymological (LDH, G6PD) analysis

**Karyology:**  $2n=42$ , variability in the range between 36-42 chromosomes, modal number of chromosomes 39, some cells have 1-2 large acrocentric chromosomes (routine dye), number of poliploid cells 1.0%

**Other properties:** synthesise several specific proteins characteristic of muscle tissue. Differentiates forming multinucleated muscle fiber

**Applications:** differentiation, cell biology

**Collections:** ATCC CRL 1769; SPBIC



**Origin:** mouse DBA/2, lymphocytic leukemia, ascitic fluid.  
J.Natl.Cancer Inst. 1966. 36: 405-421.

**Morphology:** lymphoblast-like

**Mode of cultivation:** suspension

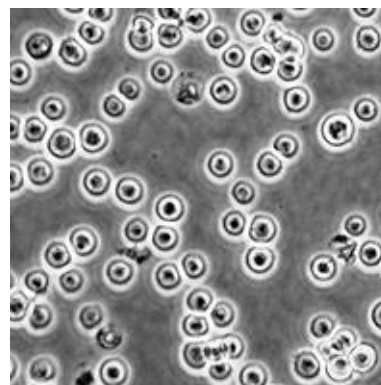
**Conditions for cultivation:** medium - RPMI 1640

serum - FBS 10%

subculture procedure - optimal

population density  $5.0 \times 10^4$  -  $8.0 \times 10^5$   
cells/ml

cryoconservation - growth medium,  
10% DMSO,  $3.0 \times 10^6$  cells/ml in  
ampule



**Viability after cryoconservation:** 90% (0 passage,  
dye trypan blue)

**Sterility:** tests for bacteria, fungi and mycoplasma were negative

**Species:** karyological and isoenzymological (LDH, G6PD) analysis

**Karyology:**  $2n=40$ , variability in the range between 34-42 chromosomes, modal  
number of chromosomes 39-41, number of polyploid cells 0.2%.

**Tumorigenicity:** tumorigenic in singeneic and nude mice

**Applications:** cytotoxicity, tumorigenicity, cell biology.

**Collections:** ATCC CCL 219; ECACC 87092804; SPBIC.

**Origin:** rat, skeletal muscle cells transformed by methylcholanthrene, derived from L6.  
Exp.Cell Res. 1979. 120: 1; Cytology (Russ). 1983, 25: 1096-1097;.

**Morphology:** myoblast-like

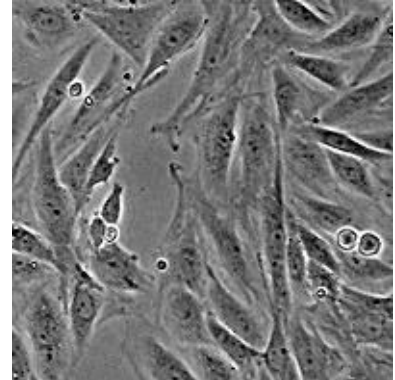
**Mode of cultivation:** monolayer

**Conditions for cultivation:** medium - DMEM

serum - FBS 10%

subculture procedure - cells detach from flask using trypsin 0.25%: EDTA 0.02% (1:3), split ratio 1:4 - 1:6, do not allow cultures to become completely confluent.

cryoconservation - growth medium, 10% DMSO,  $1.0 \times 10^6$  cells/ml in ampule



**Viability after cryoconservation:** 85% (0 passage, dye trypan blue)

**Sterility:** tests for bacteria, fungi and mycoplasma were negative

**Species:** karyological, isoenzymological (LDH, G6PD) and immunofluorescent analysis

**Karyology:**  $2n=42$ , variability in the range between 41-47 chromosomes, modal number of chromosomes 42-43, number of markers- 3-5 (differential dye) some cells have one small submetacentric chromosome with gap in short arm microchromosoma (routine dye), number of polyploid cells 5%.

**Plating efficiency:** 42%

**Other properties:** differentiates producing myotubes, synthesis of muscle specific proteins.

**Applications:** differentiation, myogenesis.

**Collections:** SPBIC.

## LLC-MK2, derivative

**Origin:** rhesus monkey, kidney, derived from LLC-MK2 original.  
Anat.Res. 1956. 124: 490; J.Gen.Virol. 1979. 43: 289.

**Morphology:** epithelial-like

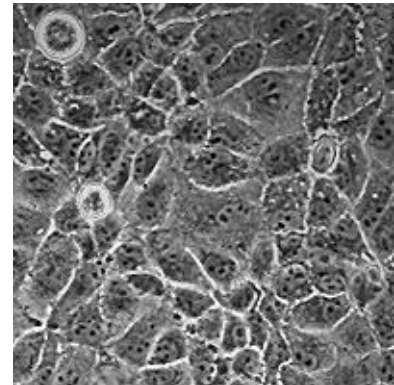
**Mode of cultivation:** monolayer

**Conditions for cultivation:** medium - DMEM

serum - FBS 10%

subculture procedure - cells detach from flask using trypsin 0.25%: EDTA 0.02% (1:3), split ratio 1:3, optimal population density  $2.0-4.0 \times 10^4$  cells/cm<sup>2</sup>

cryoconservation - growth medium, 5 – 7%DMSO,  $1.0-1.5 \times 10^6$  cells/ml in ampule



**Viability after cryoconservation:** 80-90% (0 passage, dye trypan blue)

**Sterility:** tests for bacteria, fungi and mycoplasma were negative

**Species:** karyological and isoenzymological (LDH, G6PD and nucleoside phosphorylase) analysis

**Karyology:**  $2n=42$ , variability in the range between 63-73 chromosomes, modal number of chromosomes 67-70, number of markers - 1-4 middle submetacentrics with the second constriction, number of polyploid cells 4.8%.

**Plating efficiency:** 45%.

**Other properties:** virus susceptibility: poliovirus 1, 2, 3, parainfluenza 2, 3

**Applications:** virology.

**Collections:** ATCC CCL 7.1; SPBIC; SPBII.

L-M (TK<sup>-</sup>, APRT<sup>-</sup>)

**Origin:** mouse, connective, derived from NCTC clone 929.

Submitted Institute of Biochemistry, Martinsried, FGR.

**Morphology:** fibroblast-like

**Mode of cultivation:** monolayer

**Conditions for cultivation:** medium - DMEM

serum - FBS 10%

subculture procedure - cells detach

from flask using trypsin 0.25%: EDTA

0.02% (1:3), split ratio 1:3 - 1:5,

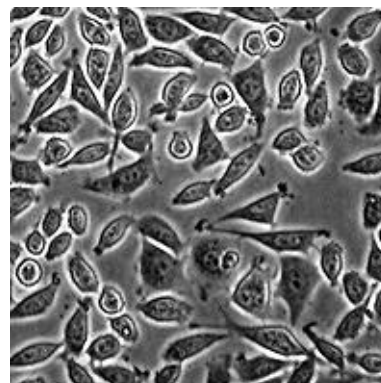
optimal population density  $2.0-3.0 \times 10^4$

cells/cm<sup>2</sup>

cryoconservation - growth medium,

10% FBS, 5-10% DMSO,  $1.8 \times 10^6$

cells/ml in ampule



**Viability after cryoconservation:** 68% (0 passage, dye trypan blue)

**Sterility:** tests for bacteria, fungi and mycoplasma were negative

**Species:** karyological, isoenzymological (LDH, G6PD) and immunological analysis

**Karyology:**  $2n=40$ , variability in the range between 46-51 chromosomes, modal number of chromosomes 49, number of markers - 9 metacentrics (routine dye).

**Plating efficiency:** 25%

**Tumorigenicity:** non tumorigenic

**Other properties:** deficient in thymidine kinase and adenine phosphoribosyl transferase (resistant to 5-bromodeoxyuridine and 8-azaadenine.

Retrovirus type A production

**Applications:** virology, somatic cell genetics, cell biology.

**Collections:** SPBIC.



**Origin:** mouse C3H/An, connective, derived from NCTC clone 929.  
Proc.Roy.Soc. 1967. 168: 431-438.

**Morphology:** round cells

**Mode of cultivation:** suspension

**Conditions for cultivation:** medium - EMEM

serum - FBS 10%

other components - NEAA 1%

subculture procedure - optimal

population density  $0.8-1.0 \times 10^5$  cells/ml

cryoconservation - growth medium,

10% DMSO,  $2.0 \times 10^6$  cells/ml in ampule

**Viability after cryoconservation:** 87% (0 passage,  
dye trypan blue)

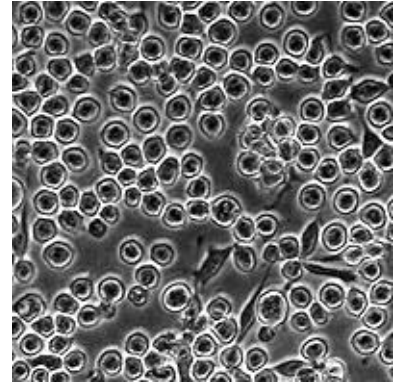
**Sterility:** tests for bacteria, fungi and mycoplasma were negative

**Species:** karyological, isoenzymological (LDH, G6PD) and immunofluorescent analysis

**Karyology:**  $2n=40$ , variability in the range between 53-57 chromosomes, modal  
number of chromosomes 55-56, number of polyploid cells 1%.

**Applications:** biochemistry, cell biology.

**Collections:** SPBIC.



**Origin:** mouse, connective, LS cells adapted to monolayer growth

Tsitologiya 1981.23.10.1216

**Morphology:** fibroblast-like

**Mode of cultivation:** monolayer

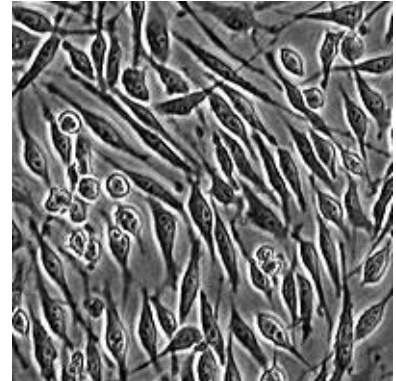
**Conditions for cultivation:** medium - EMEM

serum - FBS 10%

other components - NEAA 1%

subculture procedure - cells detach from flask using trypsin 0.25%: EDTA 0.02% (1:3), split ratio 1:4, optimal population density  $2.0-4.0 \times 10^4$  cells/cm<sup>2</sup>

cryoconservation - growth medium, 10% DMSO,  $1.0 \times 10^6$  cells/ml in ampule



**Viability after cryoconservation:** 92% (0 passage, dye trypan blue)

**Sterility:** tests for bacteria, fungi and mycoplasma were negative

**Species:** karyological, isoenzymological (LDH, G6PD) and immunofluorescent analysis

**Karyology:**  $2n=40$ , variability in the range between 52-58 chromosomes, modal number of chromosomes 56, most cells have 1 metacentric with second constriction (routine dye), number of polyploid cells 2%.

**Tumorigenicity:** tumorigenic in syngenic animals

**Applications:** oncology, biochemistry.

**Collections:** SPBIC.

## McCoy B

**Origin:** mouse, cells obtained from synovial fluid of human knee joint with arthritis (Z. Zellforsch. 1957, 47: 158), but later one of sublines proved to be of mouse origin. Proc. Soc. Exp. Biol. Med. 1965, 118: 354.

**Morphology:** fibroblast-like

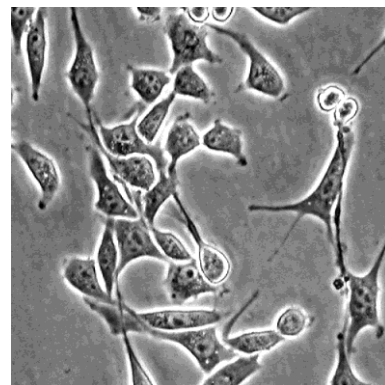
**Mode of cultivation:** monolayer

**Conditions for cultivation:** medium - DMEM

serum - FBS 10%

subculture procedure - cells detachment using EDTA 0.04 %, split ratio 1:3 - 1:7

cryoconservation - growth medium, 5% DMSO,  $1.0-1.5 \times 10^6$  cells/ml in ampule.



**Viability after cryoconservation:** 80 - 90 % (0 passage, dye trypan blue)

**Sterility:** tests for bacteria, fungi and mycoplasma were negative

**Species:** karyological and isoenzymological (LDH, G6PD) analysis

**Karyology:**  $2n=40$ , variability in the range between 55-63 chromosomes, modal number of chromosomes 58-60, number of markers - 1 small telocentric chromosome, some cells have dicentric chromosomes (routine dye), number of polyploid cells 2.6%.

**Other properties:** virus susceptibility: vesicular stomatitis.

Susceptibility to chlamidia.

**Applications:** cell biology, virology

**Collections:** ATCC CRL 1696, ECACC 90010305, SPBII, SPBIC.

**Origin:** mouse C3H, rhabdomyosarcoma induced by methylcholanthrene.  
Cytology, Russ. 1970. 12: 798.

**Morphology:** fibroblast-like

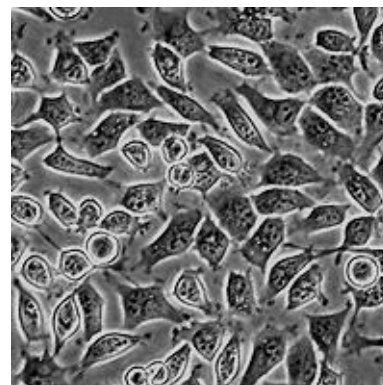
**Mode of cultivation:** monolayer

**Conditions for cultivation:** medium - DMEM

serum - FBS 10%

subculture procedure - cells detach  
from flask using trypsin 0.25%: EDTA  
0.02% (1:3), split ratio 1:3 - 1:8

cryoconservation - growth medium,  
10% DMSO,  $1.0 \times 10^6$  cells/ml in  
ampule



**Viability after cryoconservation:** 91% (0 passage, dye trypan blue)

**Sterility:** tests for bacteria, fungi and mycoplasma were negative

**Species:** karyological and isoenzymological (LDH, G6PD) analysis

**Karyology:**  $2n = 40$ , variability in the range between 57-85 chromosomes, 77-78 chromosomes in 30% of cells, some cells have 1-3 microchromosomes.

**Plating efficiency:** 80%

**Tumorigenicity:** tumorigenic in syngeneic animals

**Applications:** tumorigenicity

**Collections:** SPBIC.

**Origin:** mouse DBA/2, rhabdomyosarcoma induced by methylcholanthrene.  
Cytology, Russ. 1988. 30: 726.

**Morphology:** fibroblast-like

**Mode of cultivation:** monolayer

**Conditions for cultivation:** medium - DMEM

serum - FBS 10 %

subculture procedure - cells detach  
from flask using trypsin 0.25%: EDTA  
0.02% (1:3), split ratio 1:3 - 1:8

cryoconservation - growth medium,  
10% DMSO,  $1.0 \times 10^6$  cells/ml in ampule

**Viability after cryoconservation:** 90% (0 passage,  
dye trypan blue)

**Sterility:** tests for bacteria, fungi and mycoplasma were negative

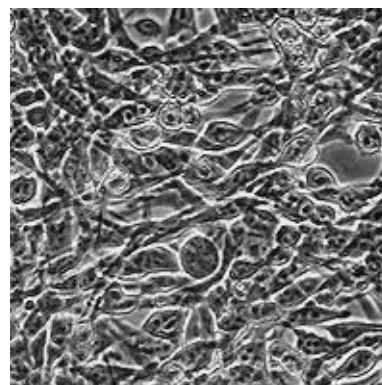
**Species:** karyological and isoenzymological (LDH, G6PD) analysis

**Karyology:**  $2n=40$ , variability in the range between 50-60 chromosomes, modal  
number of chromosomes 53, number of markers - 2 (differential dye)

**Tumorigenicity:** tumorigenic in syngeneic animals

**Applications:** tumorigenicity:

**Collections:** SPBIC.



## MDBK (NBL-1)

**Origin:** bovine, kidney.

Proc.Soc.Exp.Biol.Med.1958. 98:574; J.Natl.Cancer Inst.1986. 76:87-93.

**Morphology:** epithelial-like

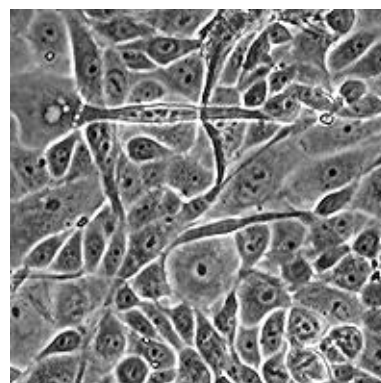
**Mode of cultivation:** monolayer

**Conditions for cultivation:** medium - F12

serum - FBS 10%

subculture procedure - cells detach from flask using trypsin 0.25%: EDTA 0.02% (1:3), split ratio 1:3 - 1:5, optimal population density  $2.0-4.0 \times 10^4$  cells/cm<sup>2</sup>

cryoconservation - growth medium, 10% DMSO,  $1.0 \times 10^6$  cells/ml in ampule



**Viability after cryoconservation:** 90% (0 passage, dye trypan blue)

**Sterility:** tests for bacteria, fungi and mycoplasma were negative

**Species:** karyological and isoenzymological (LDH, G6PD) analysis

**Karyology:**  $2n=60$ , variability in the range between 40-57 chromosomes, modal number of chromosomes 51-53, number of markers - 11-14 (differential dye), number of polyploid cells 2.0%.

**Plating efficiency:** 19%.

**Other properties:** virus susceptibility: - alphaviruses, vesicular stomatitis, IBR, BVD, bovine parvoviruses, bovine adenoviruses I and III, parainfluenza 3.

**Applications:** virology.

**Collections:** ATCC CCL 22; ECACC 90050801; SPBIC; MWIIW.

## MDCK (NBL-2)

**Origin:** dog, kidney.

Proc.Soc.Exp.Biol.Med. 1958. 98: 574.

**Morphology:** epithelial-like

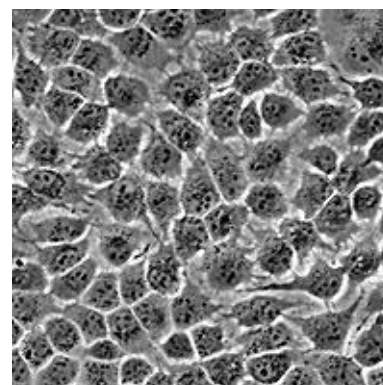
**Mode of cultivation:** monolayer

**Conditions for cultivation:** medium - DMEM

serum - FBS 10%

subculture procedure - cells detach from flask using trypsin 0.25%: EDTA 0.02% (1:3), split ratio 1:3 - 1:6, optimal population density  $1.0-3.0 \times 10^4$  cells/cm<sup>2</sup>

cryoconservation - growth medium, 10% DMSO,  $1.0 \times 10^6$  cells/ml in ampule



**Viability after cryoconservation:** 90% (0 passage, dye trypan blue)

**Sterility:** tests for bacteria, fungi and mycoplasma were negative

**Species:** karyological analysis

**Karyology:**  $2n=78$ , variability in the range between 75-83 chromosomes, modal number of chromosomes 78-80, number of markers - 1-2 large submetacentric chromosomes, some cells have 1-2 middle meta- or submetacentric chromosomes (routine dye), number of polyploid cells 0.6%.

**Plating efficiency:** 35%.

**Other properties:** virus susceptibility: vesicular stomatitis, vaccinia, Coxsackie B-5, reovirus 2, 3; adenovirus 4, 5; influenza A, B, C; carnivorous plague, arboviruses, arenaviruses, infectious canine hepatitis, swine vesicular exanthema.

**Applications:** virology, biotechnology, cell biology.

**Collections:** ATCC CCL 34; ECACC 84121903; 85011435; MWIIV; ESCC; SPBIC.

**Origin:** chicken, lymphoblastoma.

Submitted from Fridrich Loeffler Institute, Germany.

**Morphology:** lymphoblast-like

**Mode of cultivation:** suspension

**Conditions for cultivation:** medium - EMEM

serum - FBS 10%

other components - NEAA 1%

subculture procedure - optimal

population density  $2.0 \times 10^5$  cells/cm<sup>2</sup>

cryoconservation - growth medium,

10% DMSO,  $5.0-6.0 \times 10^6$  cells/ml in

ampule

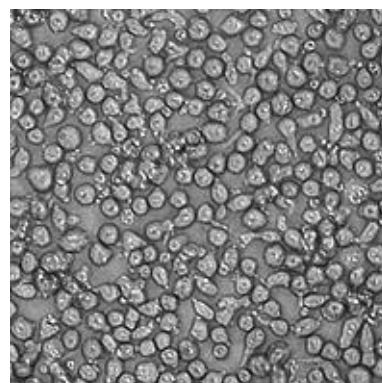
**Viability after cryoconservation:** 96% (0 passage, dye trypan blue)

**Sterility:** tests for bacteria, fungi and mycoplasma were negative

**Species:** karyological analysis

**Applications:** cell biology

**Collections:** SPBIC.





**Origin:** mouse C3HA, hepatoma.

Bull.Exp.Biol.Med. Russ. 1972. 5: 94-95; Atlas of chromosomes of human and animal cell lines, S.E. Mamaeva, 2002. Moscow, Scientific world

**Morphology:** epithelial-like

**Mode of cultivation:** monolayer

**Conditions for cultivation:** medium - DMEM

serum - FBS 10%

subculture procedure - cells detach from flask using trypsin 0.25%: EDTA 0.02% (1:3), split ratio 1:3 - 1:6

cryoconservation - growth medium, 10% DMSO,  $1.0 \times 10^6$  cells/ml in ampule

**Viability after cryoconservation:** 98% (0 passage, dye trypan blue)

**Sterility:** tests for bacteria, fungi and mycoplasma were negative

**Species:** karyological, isoenzymological (LDH and G6PD) and immunofluorescent analysis

**Karyology:**  $2n=40$ , variability in the range between 50-60 chromosomes, modal number of chromosomes 55, number of markers - 2 large and middle submetacentric chromosomes, some cells have middle telocentric chromosome with secondary constriction (routine dye).

**Tumorigenicity:** tumorigenic in syngeneic animals

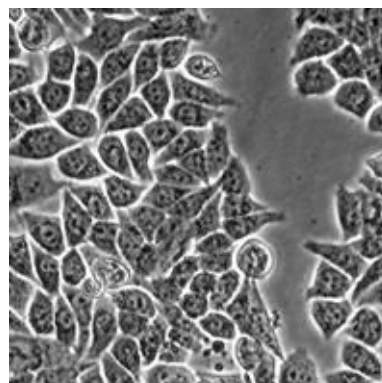
**Other properties:**

virus susceptibility: adenovirus 6.

Transferrin synthesis

**Applications:** tumorigenicity, cell biology.

**Collections:** SPBIC.



**Origin:** mink, lung.

Virology 1974. 60: 282-287.

**Morphology:** epithelial-like

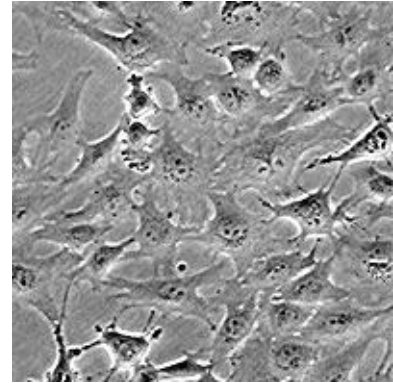
**Mode of cultivation:** monolayer

**Conditions for cultivation:** medium - DMEM

serum - FBS 10%

subculture procedure - cells detach from flask using trypsin 0.25%: EDTA 0.02% (1:3), split ratio 1:3 - 1:6, optimal population density  $2.0-4.0 \times 10^4$  cells/cm<sup>2</sup>

cryoconservation - growth medium, 10% DMSO,  $1.0 \times 10^6$  cells/ml in ampule



**Viability after cryoconservation:** 90% (0 passage, dye trypan blue)

**Sterility:** tests for bacteria, fungi and mycoplasma were negative

**Species:** karyological analysis

**Karyology:**  $2n=30$ , variability in the range between 24-32 chromosomes, modal number of chromosomes 30, pseudodiploid, number of markers - 1 dicentric in some cells (routine dye).

**Plating efficiency:** 5%.

**Other properties:** virus susceptibility: herpes simplex; reovirus 3; vesicular stomatitis; vaccinia; pseudorabies; IBR; murine sarcoma virus, feline sarcoma virus.

**Applications:** virology.

**Collections:** ATCC CCL 64; ECACC 88050503; MWIIV; SPBIC.

**Origin:** mouse A, neuroblastoma, clone of C1300.

Proc.Natl.Acad.Sci. 1962. 48: 1184-1190.

**Morphology:** neuroblast-like

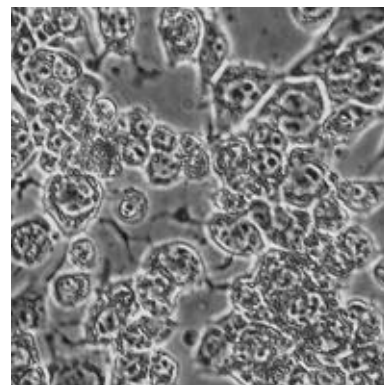
**Mode of cultivation:** monolayer

**Conditions for cultivation:** medium - F10

serum - HS 12.5%, FBS 2.5%

subculture procedure - cells detach from flask using trypsin 0.25%: EDTA 0.02% (1:3), split ratio 1:2, optimal population density  $3.0-5.0 \times 10^4$  cells/cm<sup>2</sup>

cryoconservation - growth medium, 10% DMSO,  $2.0 \times 10^6$  cells/ml in ampule



**Viability after cryoconservation:** 85% (0 passage, dye trypan blue)

**Sterility:** tests for bacteria, fungi and mycoplasma were negative

**Species:** karyological and immunofluorescent analysis

**Karyology:**  $2n=40$ , variability in the range between 67-99 chromosomes without modal number, number of markers - 6-10 metacentrics (routine dye).

**Plating efficiency:** 80%.

**Other properties:** virus susceptibility: vesicular stomatitis, herpes simplex, vaccinia. Acetylcholinesterase, choline acetylase and tyrosine hydroxylase production.

**Applications:** tumorigenicity, enzymology, virology, differentiation.

**Collections:** ATCC CCL 147; ECACC 89121405; SPBIC.

## NCTC clone 929

**Origin:** mouse C3H/An, connective, clone of cell line L.

J.Natl.Cancer Inst. 1943. 4: 165; J.Natl.Cancer Inst. 1948. 9: 229; J.Natl.Cancer Inst. 1951. 12: 133; 1953. 14: 655; Cancer Res. 1956. 16: 162;

J.Biophys.Biochem.Cytol.1958. 4: 567; Natl.Cancer Inst.Monogr.1962. 7: 147; Atlas of chromosomes of human and animal cell lines, S.E. Mamaeva, 2002. Moscow, Scientific world

**Morphology:** fibroblast-like

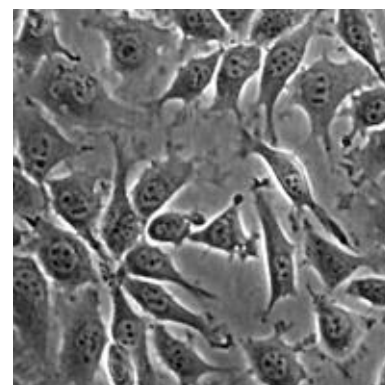
**Mode of cultivation:** monolayer

**Conditions for cultivation:** medium - DMEM

serum - FBS 10%

subculture procedure - cells detach from flask using trypsin 0.25%: EDTA 0.02% (1:3), split ratio 1:3 - 1:6, optimal population density  $1.0-3.0 \times 10^4$  cells/cm<sup>2</sup>

cryoconservation - growth medium, 5 – 10% DMSO,  $1.0-1.5 \times 10^6$  cells/ml in ampule



**Viability after cryoconservation:** 80-90% (0 passage, dye trypan blue)

**Sterility:** tests for bacteria, fungi and mycoplasma were negative

**Species:** karyological, isoenzymological (LDH and G6PD) and immunofluorescent analysis

**Karyology:**  $2n=40$ , variability in the range between 58-66 chromosomes, modal number of chromosomes 64-65, number of markers - 29 including 1 polycentric (differential dye), number of polyploid cells 1%.

**Plating efficiency:** 40%.

**Tumorigenicity:** tumorigenic in syngeneic animals

**Other properties:** virus susceptibility: pseudorabies, vesicular stomatitis, paramixovirus, togaviruses, herpes simplex.

Susceptibility to chlamidia

**Applications:** tumorigenicity, differentiation, virology, biotechnology.

**Collections:** ATCC CCL 1; ECACC 88102702; MWIIW; SPBIC; SPBII.

**Origin:** mouse A (albino), neuroblastoma.

J.Cell Biol. 1969. 43: 69A; Proc.Natl.Acad.Sci. 1970. 65: 129-136.

**Morphology:** neuron-like and amoeboid-like.

**Mode of cultivation:** monolayer

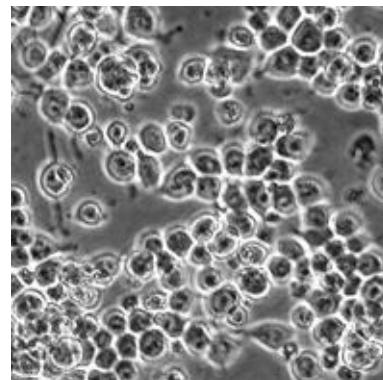
**Conditions for cultivation:** medium - EMEM

serum - FBS 10%

other components - NEAA 1%

subculture procedure - cells detach from flask using EDTA 0.02%, split ratio 1:2 - 1:4, optimal population density  $2.0-4.0 \times 10^4$  cells/cm<sup>2</sup>

cryoconservation - growth medium, 8%DMSO,  $2.0-3.0 \times 10^6$  cells/ml in ampule



**Viability after cryoconservation:** 91% (0 passage, dye trypan blue)

**Sterility:** tests for bacteria, fungi and mycoplasma were negative

**Species:** karyological, isoenzymological (LDH and G6PD) and immunofluorescent analysis

**Karyology:**  $2n=40$ , variability in the range between 70-96 chromosomes without modal number, 32% of cells have middle metacentric chromosome with gap (routine dye), each cell have 1-7 microchromosomes.

**Plating efficiency:** 60%.

**Tumorigenicity:** tumorigenic in syngeneic animals.

**Other properties:**

Virus susceptibility: vesicular stomatitis (Indiana), herpes simplex.

Microtubular protein synthesis

**Applications:** differentiation, tumorigenicity, neurophysiology, cytoskeleton study.

**Collections:** ATCC CCL 131; ECACC 89121404; SPBIC.

**Origin:** NIH/Swiss mouse, embryo.

J. Virology 1960. 4: 549-553; J. Cell Biol. 1963. 17: 299; J. Virology 1969. 4: 549-556; Science 1973. 182: 1151; Cell 1979. 16: 63-75; and 347-356.

**Morphology:** fibroblast-like

**Mode of cultivation:** monolayer

**Conditions for cultivation:** medium - DMEM

serum - FBS 10%

subculture procedure - cells detach from flask using trypsin 0.25%: EDTA 0.02% (1:3), split ratio 1:3 - 1:8, optimal

population density  $2.0-4.0 \times 10^4$  cells/cm<sup>2</sup>

cryoconservation - growth medium, 10% DMSO,  $1.0-1.5 \times 10^6$  cells/ml in ampule

**Viability after cryoconservation:** 93% (0 passage, dye trypan blue)

**Sterility:** tests for bacteria, fungi and mycoplasma were negative

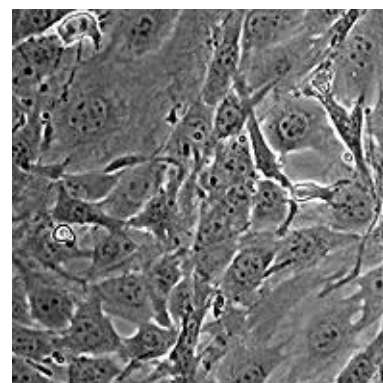
**Species:** karyological and isoenzymological (LDH, G6PD) analysis

**Karyology:**  $2n=40$ , variability in the range between 65-73 chromosomes, modal number of chromosomes 70, number of markers - 1 (routine dye), 1-2 microchromosomes in the most cells, number of polyploid cells 1.2%.

**Other properties:** virus susceptibility: vesicular stomatitis, herpes simplex, vaccinia, murine leukemia, murine sarcoma virus, N-tropic oncornaviruses C. Contact inhibition of growth (by density  $8-10 \times 10^4$  cells/cm<sup>2</sup>).

**Applications:** tumorigenicity, genetical transformation, cell biology.

**Collections:** ATCC CRL 1658; DSM ACC 59; MWIIW; SPBIC.



**Origin:** rat, kidney.

J.Cell Physiol. 1978. 94: 35-342.

**Morphology:** fibroblast-like

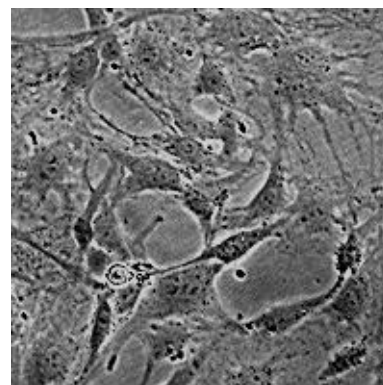
**Mode of cultivation:** monolayer

**Conditions for cultivation:** medium - F10

serum - FBS 10%

subculture procedure - cells detach from flask using trypsin 0.25%: EDTA 0.02% (1:3), split ratio 1:3 - 1:6

cryoconservation - growth medium, 10% DMSO,  $1.0-2.0 \times 10^6$  cells/ml in ampule



**Viability after cryoconservation:** 80% (0 passage, dye trypan blue)

**Sterility:** tests for bacteria, fungi and mycoplasma were negative

**Species:** karyological and isoenzymological (LDH, G6PD) analysis

**Karyology:**  $2n=42$ , variability in the range between 37-43 chromosomes, modal number of chromosomes 40, number of markers - 1 (routine dye), some cells have 1-2 dicentrics and 1-4 microchromosomes, number of polyploid cells 14%.

**Other properties** : virus susceptibility: murine sarcoma virus.

EGF receptors.

**Applications:** genetical transformation, cell biology.

**Collections:** ATCC CRL 1570; ECACC 86101301; SPBIC.

**Origin:** mouse, clone of myeloma P3X63Ag8.

Methods Enzymol. 1981. 73B: 3.

**Morphology:** lymphoblast-like

**Mode of cultivation:** suspension

**Conditions for cultivation:** medium - DMEM/F12

serum - FBS 10%

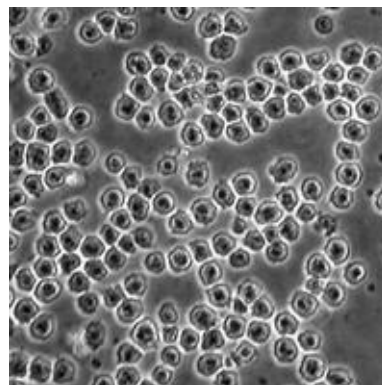
subculture procedure - optimal

population density  $5.0-9.0 \times 10^5$  cells/ml

cryoconservation - growth medium,

10% DMSO,  $5.0-6.0 \times 10^6$  cells/ml in

ampule



**Viability after cryoconservation:** 85% (0 passage, dye trypan blue)

**Sterility:** tests for bacteria, fungi and mycoplasma were negative

**Species:** karyological and isoenzymological (LDH, G6PD) analysis

**Karyology:**  $2n=40$ , variability in the range between 40-65 chromosomes, modal number of chromosomes 60, number of markers - 2-5 meta- and submetacentric chromosomes (routine dye), number of polyploid cells 2.8%.

**Other properties:** does not synthesize Ig.

Resistant to 8-azaguanine

**Applications:** fusion partner for hybridomas.

**Collections:** MWIHW; SPBIC.



### P3/NS1/1-Ag4-1(NS-1)

**Origin:** mouse BALB/c, myeloma, clone of P3X63Ag8.

Exp. Cell Res. 1970. 60:61; J. Mol. Biol. 1974. 90: 691; Eur. J. Immunol. 1976. 6: 511.

**Morphology:** lymphoid

**Mode of cultivation:** suspension

**Conditions for cultivation:** medium - RPMI 1640

serum - FBS 10%

subculture procedure optimal population density 1.0-5.0x10<sup>5</sup> cells/ml

cryoconservation - growth medium, 10% DMSO, 5.0x10<sup>6</sup> cells/ml in ampule

**Viability after cryoconservation:** 80% (0 passage, dye trypan blue)

**Sterility:** tests for bacteria, fungi and mycoplasma were negative

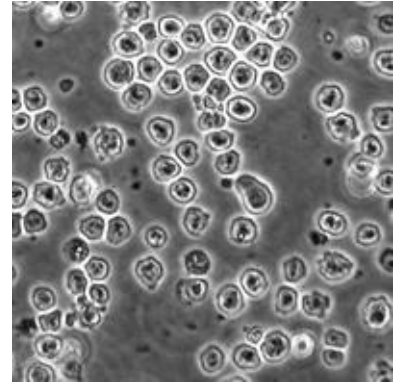
**Species:** karyological and isoenzymological (LDH, G6PD) analysis

**Other properties:** does not secrete Ig.

Resistant to 8-azaguanine

**Applications:** fusion partner for hybridomas, tumorigenicity.

**Collections:** ATCC TIB 18; DSM ACC 145; ECACC 85011427; MWIIV; SPBIC.



**Origin:** mouse BALB/c, myeloma, clone of P3X63Ag8.  
J.Immunol. 1979. 123: 1548.

**Morphology:** lymphoid

**Mode of cultivation:** suspension

**Conditions for cultivation:** medium - RPMI 1640

serum - FBS 10

subculture procedure optimal population  
density  $3.0-5.0 \times 10^5$  cells/ml

cryoconservation - growth medium,  
10% DMSO,  $5.0-6.0 \times 10^6$  cells/ml in  
ampule

**Viability after cryoconservation:** 71% (0 passage, dye  
trypan blue)

**Sterility:** tests for bacteria, fungi and mycoplasma were negative

**Species:** karyological and isoenzymological (LDH and G6PD) analysis

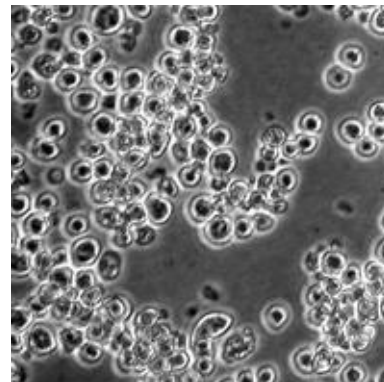
**Karyology:**  $2n=40$ , variability in the range between 46-61 chromosomes, modal  
number of chromosomes 51-53, number of markers - 1-3 meta- and submetacentric  
chromosomes (routine dye), number of polyploid cells 2%.

**Other properties:** does not secrete Ig.

Resistant to 8- azaguanine

**Applications:** fusion partner for hybridomas, tumorigenicity.

**Collections:** ATCC CRL 1580; ECACC 85011420; DSM ACC 43; MWIIV; SPBIC.



**Origin:** mouse C3H/He, teratocarcinoma.

Dev. Biol. 1982. 89: 503-508; J. Cell Biol. 1982. 94: 253-262; Nature 1982. 299: 165-167.

**Morphology:** epithelial-like

**Mode of cultivation:** monolayer

**Conditions for cultivation:** medium -  $\alpha$ MEM

serum - FBS 10%

subculture procedure - cells detach from flask using trypsin 0.25%: EDTA 0.02% (1:3), split ratio 1:3 - 1:6

cryoconservation - growth medium, 5% DMSO,  $1.0 \times 10^6$  cells/ml in ampule

**Viability after cryoconservation:** 75% (0 passage, dye trypan blue)

**Sterility:** tests for bacteria, fungi and mycoplasma were negative

**Species:** karyological and isoenzymological (LDH, G6PD) analysis

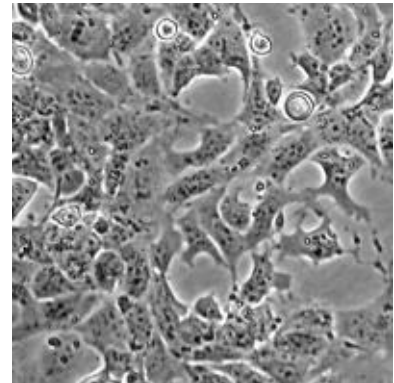
**Karyology:**  $2n=40$ , normal mouse karyotype (40, XY).

**Plating efficiency:** high efficiency in medium containing  $10^{-4}$ M  $\beta$ -mercaptoethanol.

**Other properties:** can be induced to differentiate into neuronal and glial cells in the presence of retinoic acid; in the presence of DMSO differentiate into cardiac and skeletal muscle.

**Applications:** differentiation.

**Collections:** ATCC CRL 1825; SPBIC.



**Origin:** mouse DBA/2, lymphoid neoplasm induced by methylcholanthrene.

Am.J.Pathol. 1957. 33: 603.

**Morphology:** lymphoblast-like

**Mode of cultivation:** suspension

**Conditions for cultivation:** medium - RPMI 1640

serum - FBS 10%

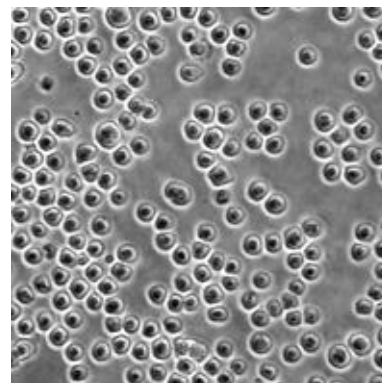
subculture procedure - optimal

population density  $1.0 \times 10^5$  cells/ml

cryoconservation - growth medium,

10% DMSO,  $3.0-4.0 \times 10^6$  cells/ml in

ampule



**Viability after cryoconservation:** 80% (0 passage, dye trypan blue)

**Sterility:** tests for bacteria, fungi and mycoplasma were negative

**Species:** karyological and isoenzymological (LDH, G6PD) analysis

**Karyology:**  $2n = 40$ , variability in the range between 38-44 chromosomes, modal number of chromosomes 41-42, number of markers - 6 (differential dye), the most cells have 3-5 microchromosomes including double minute chromosomes, number of polyploid cells 4.5%.

**Plating efficiency:** the cells cannot be plated.

**Tumorigenicity:** tumorigenic in nude mice

**Applications:** cell biology, tumorigenicity.

**Collections:** ATCC CCL 46; SPBIC.

**Origin:** mouse DBA/2, mastocytoma induced by methylcholanthrene. J.Natl.Cancer Inst. 1957. 18: 587; Cell Immunol. 1973. 9: 60; J.Immunol. 1973. 111: 389; J.Immunol. 1977. 119: 950; Nature 1974. 249: 49; Biochem.Biophys.Res.Commun. 1974. 61: 1268; Cancer Res. 1977. 37: 546.

**Morphology:** round cells

**Mode of cultivation:** suspension

**Conditions for cultivation:** medium - DMEM

serum - FBS 10%

subculture procedure - optimal

population density  $3.0-9.0 \times 10^5$  cells/ml

cryoconservation - growth medium, 10%

DMSO,  $5.0 \times 10^6$  cells/ml in ampule

**Viability after cryoconservation:** 70% (0 passage, dye trypan blue)

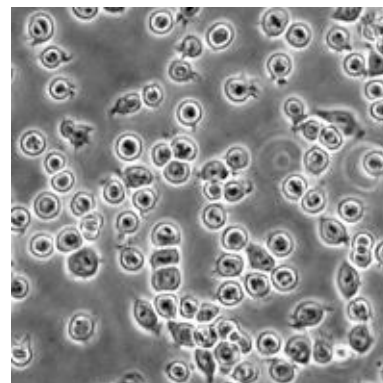
**Sterility:** tests for bacteria, fungi and mycoplasma were negative

**Species:** karyological and isoenzymological (LDH, G6PD) analysis

**Other properties:** lysozyme synthesis

**Applications:** target cell for cytotoxic T-cell assays, immunology, cell biology.

**Collections:** ATCC TIB 64; DSM ACC1; SPBIC.



**Origin:** mouse, embryo. This line was derived from NIH/3T3 TK<sup>-</sup> cells by cotransfection with retrovirus packaging construct DNA (pPAM3) and the herpes simplex virus thymidine kinase (TK) gene.

Mol.Cell Biol. 1986. 6: 2895-2902; N.Engl.J.Med. 1990. 232: 570-578.

**Morphology:** fibroblast-like

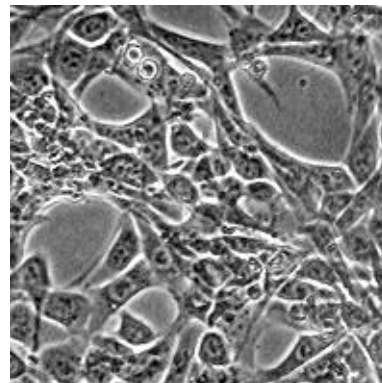
**Mode of cultivation:** monolayer

**Conditions for cultivation:** medium - DMEM

serum - FBS 10%

subculture procedure - cells detach from flask using trypsin 0.25%: EDTA 0.02% (1:3), split ratio 1:2 - 1:4, optimal population density  $3.0-5.0 \times 10^4$  cells/cm<sup>2</sup>

cryoconservation - growth medium, 10% DMSO,  $1.0 \times 10^6$  cells/ml in ampule



**Viability after cryoconservation:** 90% (0 passage, dye trypan blue)

**Sterility:** tests for bacteria, fungi and mycoplasma were negative

**Species:** karyological and isoenzymological (LDH, G6PD) analysis

**Applications:** genetical transformation, virology.

**Collections:** ATCC CRL 9078; ECACC 89032007; SPBIC.

**Origin:** pig, kidney.

Am.J.Vet.Res. 1968. 29: 153; J.Genet.Virol. 1971. 10; 195-198; Vet.microbiol. 1982. 7: 515.

**Morphology:** epithelial-like

**Mode of cultivation:** monolayer

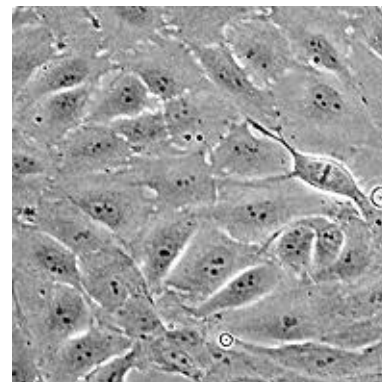
**Conditions for cultivation:** medium - EMEM

serum - FBS 10%

other components - NEAA 1%

subculture procedure - cells detach from flask using trypsin 0.25%: EDTA 0.02% (1:3), split ratio 1:5, optimal population density  $2.0-4.0 \times 10^4$  cells/cm<sup>2</sup>

cryoconservation - growth medium, 10% DMSO,  $1.0 \times 10^6$  cells/ml in ampule



**Viability after cryoconservation:** 85% (0 passage, dye trypan blue)

**Sterility:** tests for bacteria, fungi and mycoplasma were negative

**Species:** karyological and isoenzymological (LDH, G6PD) analysis

**Karyology:**  $2n=38$ , variability in the range between 30-38 chromosomes, modal number of chromosomes 37, number of markers - 1 (routine dye), number of polyploid cells 5.0%.

**Plating efficiency:** 2% (ATCC)

**Other properties:** virus susceptibility: vesicular stomatitis (Indiana); vaccinia; reovirus 2, 3; adenovirus 4, 5; Coxsackie B-2, B-3, B-4, B-5, B-6; pseudorabies; swine fever virus, swine pestis virus

**Applications:** virology.

**Collections:** ATCC CCL 33; ECACC 85022110; SPBIC.

**Origin:** mouse NIH/Swiss, embryo.

Proc.Natl.Acad.Sci. 1987. 84: 156-160; Nature 1987. 328: 131-136.

**Morphology:** fibroblast-like

**Mode of cultivation:** monolayer

**Conditions for cultivation:** medium - DMEM

serum - FBS 10%

subculture procedure - cells detach from flask using trypsin 0.25%: EDTA 0.02% (1:3), split ratio 1:4

cryoconservation - growth medium, 10% DMSO,  $1.0 \times 10^6$  cells/ml in ampule

**Viability after cryoconservation:** 97% (0 passage, dye trypan blue)

**Sterility:** tests for bacteria, fungi and mycoplasma were negative

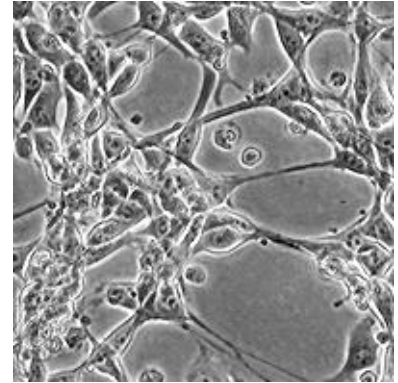
**Species:** karyological and isoenzymological (LDH, G6PD) analysis

**Karyology:**  $2n=40$ , variability in the range between 63-74 chromosomes, modal number of chromosomes 70, number of markers - 1 telocentric chromosome with secondary constriction (routine dye), 1 microchromosome, number of polyploid cells 1.5%.

**Other properties:** this line produces a vector (BAG) that can infect mouse and rat and transduce the bacterial  $\beta$  galactosidase gene.

**Applications:** genetical transformation.

**Collections:** ATCC CRL 9560; SPBIC.





**Origin:** rat kangaroo, kidney.

Nature 1962. 194: 406; Cytogenetics 1964. 3: 19.; Cytology (Russ) 1988.30: 732-738;  
Cytology (Russ) 1996. 38: 75-84

**Morphology:** epithelial-like

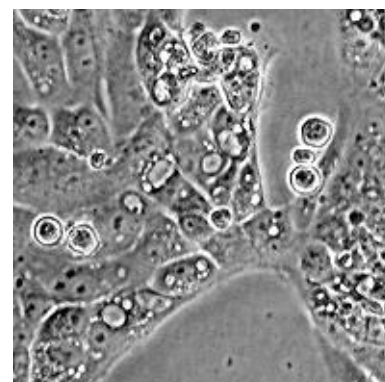
**Mode of cultivation:** monolayer

**Conditions for cultivation:** medium - DMEM

serum - FBS 10%

subculture procedure - cells detach  
from flask using trypsin 0.25%: EDTA  
0.02% (1:3), split ratio 1:2 - 1:3,  
optimal population density  $4.0-5.0 \times 10^4$   
cells/cm<sup>2</sup>

cryoconservation - growth medium,  
10% DMSO,  $1.0 \times 10^6$  cells/ml in ampule



**Viability after cryoconservation:** 98% (0 passage, dye trypan blue)

**Sterility:** tests for bacteria, fungi and mycoplasma were negative

**Species:** karyological and isoenzymological (LDH and G6PD) analysis

**Karyology:**  $2n=12$ , variability in the range between 10-17 chromosomes, modal  
number of chromosomes 11 without markers, one small metacentric of the diploid  
female karyotype is absent, number of polyploid cells 2%.

**Plating efficiency:** 2%.

**Other properties:** virus susceptibility: vesicular stomatitis (Indiana)

**Applications:** cell biology, cytogenetics, virology.

**Collections:** ATCC CCL 35; ECACC 91013163; MWIIW; SPBIC.

## PTK1 (NBL-3-17)

**Origin:** rat kangaroo, kidney, subline of Pt K1 (NBL-3)

Cytology (Russ.)1988. 30: 732-738; Cytology (Russ.) 1996. 38: 75-84; Atlas of chromosomes of human and animal cell lines, S.E. Mamaeva, 2002. Moscow, Scientific world.

**Morphology:** epithelial-like

**Mode of cultivation:** monolayer

**Conditions for cultivation:** medium - DMEM

serum - FBS 10%

subculture procedure - cells detach from flask using trypsin 0.25%: EDTA 0.02% (1:3), split ratio 1:2 - 1:3

cryoconservation - growth medium, 10% DMSO,  $1.0 \times 10^6$  cells/ml in ampule

**Viability after cryoconservation:** 88% (0 passage, dye trypan blue)

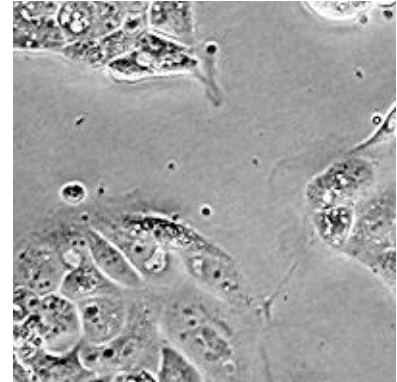
**Sterility:** tests for bacteria, fungi and mycoplasma were negative

**Species:** karyological analysis

**Karyology:**  $2n = 12$ , variability in the range between 15-19 chromosomes, modal number of chromosomes 17 without markers, hypotriploid, one small metacentric of the triploid female karyotype is absent, number of polyploid cells 3%.

**Applications:** cell biology, cytogenetics.

**Collections:** SPBIC.



**Origin:** rat, leukemic basophilic granulocyte.

Nature New Biol. 1973. 244: 73 – 76; J.Exp.Med. 1974. 139: 600 – 616.

**Morphology:** lymphoblast-like

**Mode of cultivation:** semisuspension

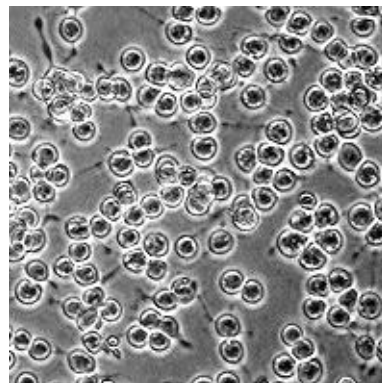
**Conditions for cultivation:** medium - EMEM

serum - FBS 10%

other components - NEAA 1%

subculture procedure - cells detach without enzymatic treatment by light shaking of flask, split ratio 1:5

cryoconservation - growth medium, DMSO 5 – 10%,  $1.0-3.0 \times 10^6$  cells/ml in ampule



**Viability after cryoconservation:** 90% (0 passage, dye trypan blue)

**Sterility:** tests for bacteria, fungi and mycoplasma were negative

**Species:** karyological analysis

**Karyology:**  $2n=42$ , variability in the range between 52-75 chromosomes, modal number of chromosomes 71-74, number of polyploid cells 0.2%.

**Other properties:** expression of FcERI (Fc of IgE); secretion of hystamin.

**Applications:** cell biology, differentiation.

**Collections:** ATCC CRL 1378; ECACC 86061001; SPBIC.

**Origin:** rat, leukemia basophilic chemically induced, peripheral blood.  
Nature New Biol. 1973. 244: 73 – 76; J.Exp.Med. 1974. 139: 600 – 616.

**Morphology:** fibroblast-like

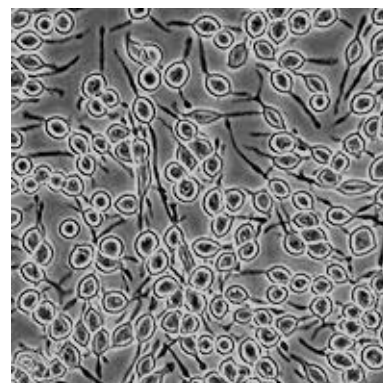
**Mode of cultivation:** monolayer

**Conditions for cultivation:** medium - EMEM

serum - FBS 15% (heat inactivated – ATCC).

subculture procedure - cells detach from flask using trypsin 0.25%: EDTA 0.02% (1:3), split ratio 1:4 - 1:8

cryoconservation - growth medium, 5 – 8% DMSO,  $2.0 \times 10^6$  cells/ml in ampule



**Viability after cryoconservation:** 80% (0 passage, dye trypan blue)

**Sterility:** tests for bacteria, fungi and mycoplasma were negative

**Species:** karyological analysis

**Other properties:** expression of FcERI (Fc of IgE);

secretion of histamin;

the cells capable to degranulation (as distinct from cell line RBL-1), i.e. to release a number of substances, in particular, histamine, associated with immune reactions.

**Applications:** cell biology, differentiation.

**Collections:** ATCC CRL 2256<sup>tm</sup>; SPBIC.

**Origin:** rat, insulinoma (pancreatic  $\beta$ -cells)

J Biol.Chem. 1996. 271: 8307-8312.

**Morphology:** epithelial-like

**Mode of cultivation:** monolayer

**Conditions for cultivation:** medium - RPMI 1640

serum - FBS 10%

other components - NEAA 1%,

HEPES 25mM

subculture procedure - cells detach from flask using trypsin 0.25%: EDTA 0.02% (1:3), split ratio 1:3

cryoconservation - growth medium, 8–10% DMSO,  $1.0-2.0 \times 10^6$  cells/ml in ampule

**Viability after cryoconservation:** 90% (0 passage, dye trypan blue)

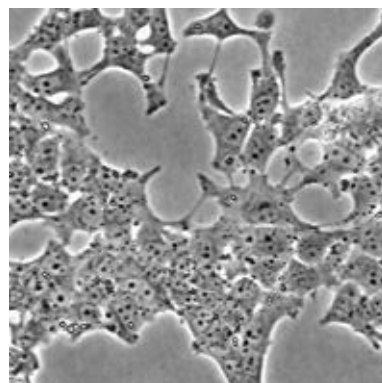
**Sterility:** tests for bacteria, fungi and mycoplasma were negative

**Species:** karyological and isoenzymological (LDH, G6PD) analysis

**Other properties:** insulin production

**Applications:** endocrinology, cell biology.

**Collections:** SPBIC.



**Origin:** rabbit, kidney.

Lancet 1963. 2: 640; J. Pathol. Bacteriol. 1968. 95: 377; Annali Sclavo 1982. 24: 336.

**Morphology:** epithelial-like

**Mode of cultivation:** monolayer

**Conditions for cultivation:** medium - EMEM

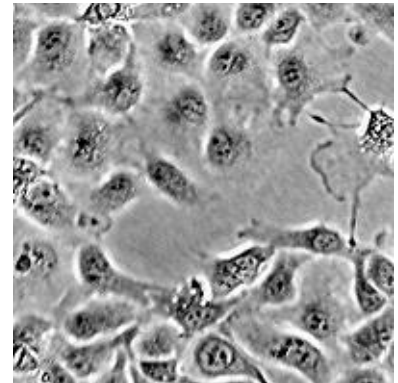
serum - FBS 10%

other components - NEAA 1%

subculture procedure - cells detach from flask using trypsin 0.25%: EDTA 0.02% (1:3), split ratio 1:2 - 1:6, optimal

population density  $2.0-4.0 \times 10^4$  cells/cm<sup>2</sup>

cryoconservation - growth medium, 5 – 10% DMSO,  $1.0-3.0 \times 10^6$  cells/ml in ampule



**Viability after cryoconservation:** 80-90% (0 passage, dye trypan blue)

**Sterility:** tests for bacteria, fungi and mycoplasma were negative

**Species:** karyological and isoenzymological (LDH and G6PD) analysis

**Karyology:**  $2n=44$ , variability in the range between 62 -68 chromosomes, modal number of chromosomes 66 , number of markers -1 large acrocentric chromosome (routine dye), number of polyploid cells 2,6%.

**Plating efficiency:** 39 %.

**Other properties:** virus susceptibility: rubella, virus B, herpes simplex, pseudorabies, vaccinia, rabbitpox, myxoma, Simian adenovirus, vesicular stomatitis, Semliki Forest virus, human enteroviruses, bovine rhynotracheitis.

**Applications:** virology.

**Collections:** ATCC CCL 37; ECACC 88062427; MWIIV; SPBII; ESCC; SPBIC.

**Origin:** rat, lymphosarcoma induced by 3,3'-dichlorobenzidine.  
Exp.Oncology (Russ.) 1980. 2: 40.

**Morphology:** lymphoblast-like

**Mode of cultivation:** suspension

**Conditions for cultivation:** medium - EMEM

serum - FBS 10%

subculture procedure - optimal

population density  $5.0-7.0 \times 10^5$  cells/ml

cryoconservation - growth medium,

10% DMSO,  $3.0-4.0 \times 10^6$  cells/ml in

ampule

**Viability after cryoconservation:** 68% (0 passage,  
dye trypan blue)

**Sterility:** tests for bacteria, fungi and mycoplasma were negative

**Species:** immunofluorescent and isoenzymological (LDH, G6PD) analysis

**Karyology:**  $2n=42$ , variability in the range between 34-58 chromosomes, modal  
number of chromosomes 38-42.

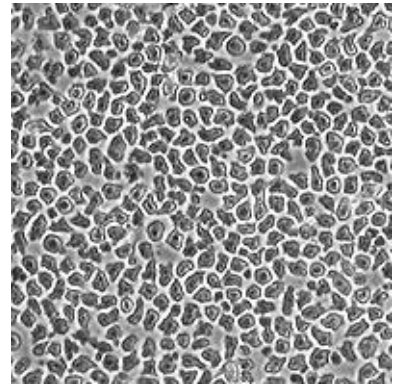
**Tumorigenicity:** tumorigenic in syngeneic animals

**Other properties:** retrovirus C production

short mitotic cycle (12 human)

**Applications:** tumorigenicity, immunology, virology.

**Collections:** SPBIC.



**Origin:** rabbit, cornea.

Science 1965. 149: 633; Proc.Soc.Exp.Biol.Med. 1966. 122: 783; Proc.Soc.Exp.Biol. Med. 1967. 125: 1271.

**Morphology:** fibroblast-like

**Mode of cultivation:** monolayer

**Conditions for cultivation:** medium - EMEM

serum - FBS 10%

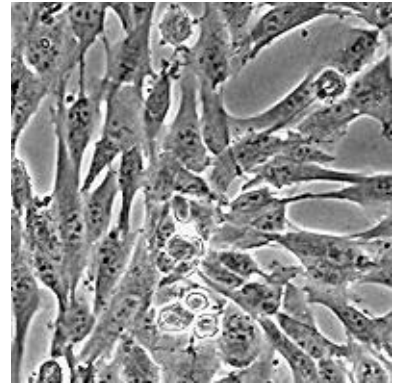
other components - NEAA 1%

subculture procedure - cells detach from flask using trypsin 0.25%: EDTA

0.02% (1:3), split ratio 1:2 - 1:4,

optimal population density 2.0-  
4.0x10<sup>4</sup> cells/cm<sup>2</sup>

cryoconservation - growth medium,  
10% DMSO, 1.0x10<sup>6</sup> cells/ml in ampule



**Viability after cryoconservation:** 85% (0 passage, dye trypan blue)

**Sterility:** tests for bacteria, fungi and mycoplasma were negative

**Karyology:** 2n= 44, variability in the range between 51-80 chromosomes, modal number of chromosomes 66, number of markers - 3-4 (routine dye), number of polyploid cells 2.5%.

**Plating efficiency:** less than 1%.

**Other properties:** virus susceptibility: rubella.

**Applications:** virology, cell biology.

**Collections:** ATCC CCL 60; ECACC 89090404; ICLC AL 96001; MWIIV; SPBIC.



## Sp2/0-Ag14

**Origin:** mouse, myeloma, hybrid of P3X63Ag8 and mouse BALB/c spleen cells.  
Nature 1978. 276: 269; J.Immunol. 1981. 126: 317-321.

**Morphology:** lymphoblast-like

**Mode of cultivation:** suspension

**Conditions for cultivation:** medium - RPMI 1640

serum - FBS 10%

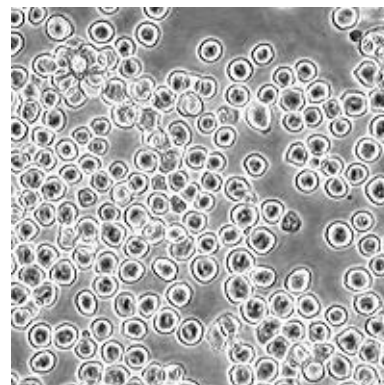
subculture procedure - optimal

population density  $3.0-9.0 \times 10^5$  cells/ml

cryoconservation - growth medium,

10% DMSO,  $5.0-6.0 \times 10^6$  cells/ml in

ampule



**Viability after cryoconservation:** 92% (0 passage, dye trypan blue)

**Sterility:** tests for bacteria, fungi and mycoplasma were negative

**Species:** karyological and isoenzymological (LDH, G6PD) analysis

**Karyology:**  $2n=40$ , variability in the range between 60-66 chromosomes, modal number of chromosomes 63-64, number of markers - 33 (differential dye).

**Plating efficiency:** 47%.

**Tumorigenicity:** tumorigenic in syngeneic animals

**Other properties:** does not secrete Ig

Resistant to 8-azaguanine.

**Applications:** fusion partner for hybridomas.

**Collections:** ATCC CRL 1581, CRL 8287; DSM ACC 146; ECACC 86072401; SPBIC.

**Origin:** pig, embryo, kidney

Abstr. 2<sup>nd</sup> Sci Conf. MNIIVP; (Russ.) 1960. 57; Atlas of chromosomes of human and animal cell lines, S.E. Mamaeva, 2002. Moscow, Scientific world

**Morphology:** epithelial-like

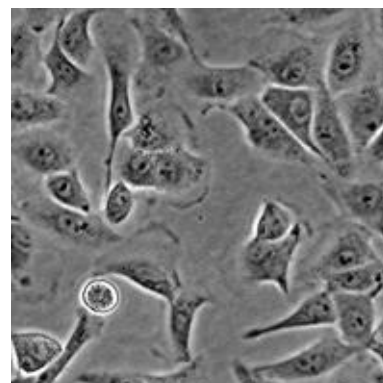
**Mode of cultivation:** monolayer

**Conditions for cultivation:** medium - DMEM

serum - FBS 10%

subculture procedure - cells detach from flask using trypsin 0.25%: EDTA 0.02% (1:3), split ratio 1:5-1:10, optimal population density  $0.9 \times 10^5$  cells/ml.

cryoconservation - growth medium, 10% DMSO,  $1.0-1.5 \times 10^6$  cells/ml in ampule



**Viability after cryoconservation:** 90-96% (0 passage, dye trypan blue)

**Sterility:** tests for bacteria, fungi and mycoplasma were negative

**Species:** karyological and isoenzymological (LDH, G6PD) analysis

**Karyology:**  $2n = 38$ , variability in the range between 39-42 chromosomes, modal number of chromosomes 40, number of markers - 1 large submetacentric chromosome (routine dye), number of polyploid cells 1,6%

**Plating efficiency:** 80%.

**Other properties:** virus susceptibility: arbovirus A and B; entero-, rota, coronaviruses of swine, rhinopneumonia of equine, influenza; encephalomyocarditis of swine, foot and mouth disease.

Presence of leukoviruses: Meson-Pfaizer-like and oncornaviruses.

**Applications:** virology, cell biology

**Collections:** MWIIW, SPBII, SPBIC, ESCC, MWIEV

**Origin:** mouse, embryonic fibroblasts, the line derived from continuous mouse line of SIM.

Proc. Natl. Acad. Sci. USA 1975. 72: 1441 – 1445; Roche Symposium on Teratomas and Differentiation, pp. 169 – 187, Sherman and Salter, eds. Academic Press, New York, 1975; Cell 1975. 6: 467 – 474; Dev. Biol. 1977. 61: 230 – 244.

**Morphology:** fibroblast-like

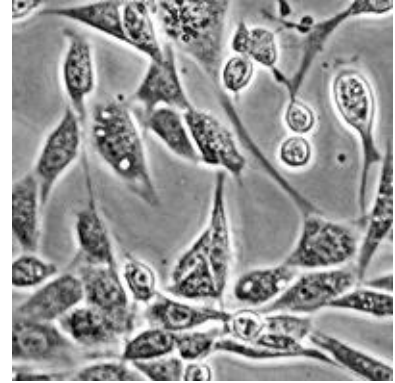
**Mode of cultivation:** monolayer

**Conditions for cultivation:** medium – DMEM

serum - FBS 10%

subculture procedure - cells detach from flask using trypsin 0.25%: EDTA 0.02% (1:3), split ratio 1:3 - 1:8, optimal population density  $2.0-4.0 \times 10^4$  cells/cm<sup>2</sup>

cryoconservation - growth medium, DMSO 5%,  $1.0-1.5 \times 10^6$  cells/ml in ampule



**Viability after cryoconservation:** 80% (0 passage, dye trypan blue)

**Sterility:** tests for bacteria, fungi and mycoplasma were negative

**Species:** karyological analysis

**Karyology:**  $2n=40$ , variability in the range between 55-65 chromosomes, modal number of chromosomes 60-62, number of markers - 2 (routine dye), 1-2 microchromosomes in the most cells, number of polyploid cells 7.0 %.

**Other properties:** resistance to 6-thioguanine and ouabain.

Sensitive to HAT medium and is HPRT negative.

**Applications:** cell biology, the cell line is used routinely to prepare feeder layer by irradiation or mitomycin C treatment in particular, for cultivation embryonic stem cells.

**Collections:** ATCC CRL 1503; ECACC 85061804; SPBIC.

**Origin:** Chinese hamster, lung

J.Cell Biol.1967.34:684; Mol. Cell Biol. 1987. 7 :4218; Atlas of chromosomes of human and animal cell lines, S.E. Mamaeva, 2002. Moscow, Scientific world

**Morphology:** fibroblast-like

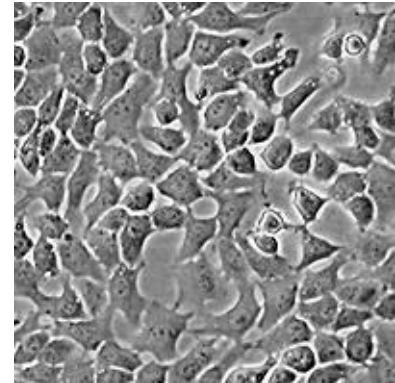
**Mode of cultivation:** monolayer

**Conditions for cultivation:** medium - DMEM

serum - FBS 10%

subculture procedure - cells detach from flask using trypsin 0.25%: EDTA 0.02% (1:3), split ratio 1:4-1:8, optimal population density  $2.0-4.0 \times 10^4$  cells/cm<sup>2</sup>

cryoconservation - growth medium, 10% DMSO,  $1.0-2.0 \times 10^6$  cells/ml in ampule



**Viability after cryoconservation:** 88 % (0 passage, dye trypan blue)

**Sterility:** tests for bacteria, fungi and mycoplasma were negative

**Species:** karyological and isoenzymological (LDH, G6PD) analysis

**Karyology:**  $2n=22$ , variability in the range between 17-23 chromosomes, modal number of chromosomes 21, number of markers 11 (differential dye), number of polyploid cells 6.0%

**Plating efficiency:** 58 %.

**Other properties:** the cells have very short G<sub>1</sub> phase of mitotic cycle

**Applications:** cell biology, proliferation mechanisms, somatic cell genetics, transformation.

**Collections:** ECACC 86041102, SPBIC.

**Origin:** African green monkey, kidney.

Nippon Rincho 1963. 21: 1209; Arch. GVS Virusforsch. 1969. 27: 379.

**Morphology:** fibroblast-like

**Mode of cultivation:** monolayer

**Conditions for cultivation:** medium - DMEM

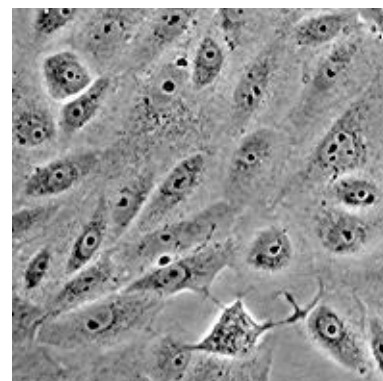
serum - FBS 10%

subculture procedure - cells detach from flask using trypsin 0.25%: EDTA 0.02% (1:2), split ratio 1:3-1:10, optimal

population density  $1.0-3.0 \times 10^4$  cells/cm<sup>2</sup>

cryoconservation - growth medium, 10%

DMSO  $1.0-2.0 \times 10^6$  cells/ml in ampule



**Viability after cryoconservation:** 77 % (0 passage, dye trypan blue)

**Sterility:** tests for bacteria, fungi and mycoplasma were negative

**Species:** karyological and isoenzymological (LDH, G6PD, nucleoside phosphorylase) analysis

**Karyology:**  $2n=60$ , variability in the range between 53-60 chromosomes, modal number of chromosomes 57-58, number of markers -3 (routine and differential dye, C banding), number of polyploid cells 2%

**Plating efficiency:** 24 %.

**Other properties:** virus susceptibility: ortomixoviruses (influenza); Getah, Ndumu, Pixuna, Ross River, Semliki, Paramaribo, Kokobera, Modoc, Murutucu, Germiston, Guaroa, Pongola, Tacaribe Arboviruses; bovine leucosis; bluetongue; adenovirus 12; paramixoviruses (parainfluenza 1 and 4, measles, respir. syncytial virus); poliovirus 3; rubella; African swine fever virus; reoviruses; herpes simplex; vesicular stomatitis; echoviruses; SV 40; SV 5.

Isoenzymes: LDG, G6PD, A, typical for primate cells.

**Applications:** virology, cell biology.

**Collections:** ATCC CCL81; ECACC 84113001, 88020401; ICLC ATL 95005; MWIIW; SPBII; ESCC; SPBIC, MWIEV.

**Origin:** African green monkey, kidney, subline of Vero.

Vero cells - Origin, properties and biomedical applications. Tokyo: Soft Science Publications. 1988. 26-29.

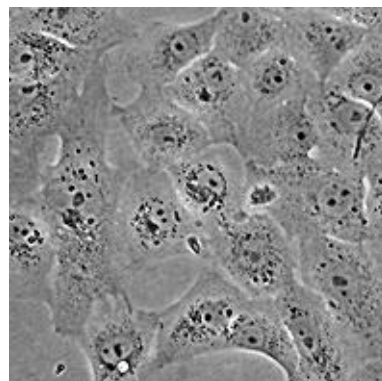
**Morphology:** fibroblast-like

**Mode of cultivation:** monolayer

**Conditions for cultivation:** medium - DMEM  
serum - FBS 10%

subculture procedure - cells detach from flask using trypsin 0.25%: EDTA 0.02% (1:3), split ratio 1:5 - 1:7, optimal population density  $1.0-3.0 \times 10^4$  cells/cm<sup>2</sup>

cryoconservation - growth medium, 10 %DMSO,  $2.0 \times 10^6$  cells/ml in ampule



**Viability after cryoconservation:** 90 % (0 passage, dye trypan blue)

**Sterility:** tests for bacteria, fungi and mycoplasma were negative

**Species:** karyological and isoenzymological (LDH, G6PD) analysis

**Karyology:**  $2n=60$ , variability in the range between 53-60 chromosomes, modal number of chromosomes 56-57, number of markers - 1 (routine dye), number of polyploid cells 9%.

**Other properties:** virus susceptibility: haemorrhagic fever viruses, Ebola.

**Applications:** virology, cell biology.

**Collections:** ATCC CRL 1587; ECACC 85020205; SPBIC.

**Origin:** mouse BALB/c, myelomonocytic leukemia.

J.Exp.Med. 1976. 143: 1528-1533; Cancer Res. 1977. 37: 546-550; J.Immunol. 1977. 119: 950-954; J.Exp.Med. 1981. 154: 1419-1431.

**Morphology:** macrophage-like

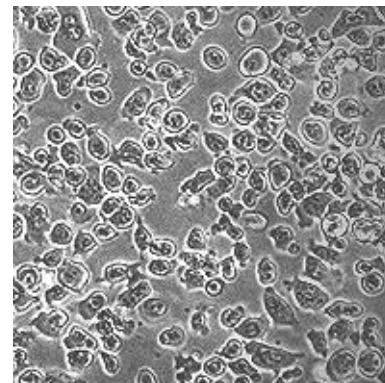
**Mode of cultivation:** semisuspension

**Conditions for cultivation:** medium - Iscove's MDM serum - FBS 10%

other components - 2-mercaptoethanol  $10^{-5}M$

subculture procedure - optimal population density  $1.0-5.0 \times 10^5$  cells/ml

cryoconservation - growth medium, 8%DMSO,  $3.0-4.0 \times 10^6$  cells/ml in ampule



**Viability after cryoconservation:** 70% (0 passage, dye trypan blue)

**Sterility:** tests for bacteria, fungi and mycoplasma were negative

**Species:** karyological and isoenzymological (LDH, G6PD) analysis

**Karyology:**  $2n=40$ , variability in the range between 72-83 chromosomes, modal number of chromosomes 75-78, number of markers - 4 metacentric chromosomes (routine dye), number of polyploid cells 0.8%.

**Other properties:** lysozyme, IL-3 and granulocyte CSA production. Ig and complement receptors.

**Applications:** immunology, cell biology, chemotherapeutic agents studies.

**Collections:** ATCC TIB 68; SPBIC.

**Origin:** mouse BALB/c, fibrosarcoma induced by methylcholathrene.

Proc.Soc.Exp.Biol.Med. 1973. 144: 813; J.Natl.Cancer Inst. 1984. 72: 23-29; Blood 1985. 65: 8-14.

**Morphology:** fibroblast-like and lymphoblast-like

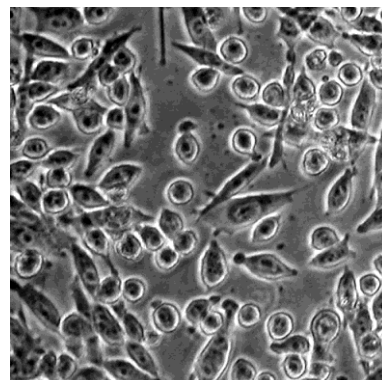
**Mode of cultivation:** monolayer

**Conditions for cultivation:** medium - RPMI 1640

serum - FBS 10%

subculture procedure - cells detach from flask using trypsin 0.25%: EDTA 0.02% (1:3), split ratio 1:2-1:3, optimal population density  $2.0-4.0 \times 10^4$  cells/cm<sup>2</sup>

cryoconservation - growth medium, 10% DMSO,  $1.0 \times 10^6$  cells/ml in ampule



**Viability after cryoconservation:** 70% (0 passage, dye trypan blue)

**Sterility:** tests for bacteria, fungi and mycoplasma were negative

**Species:** karyological and isoenzymological (LDH, G6PD) analysis

**Other properties:** the line is highly sensitive, after pretreatment with actinomycin D, to human cytotoxic monocytes, to human TNF and to lymphotoxin.

**Applications:** cytotoxicity, tumorigenicity, cell biology.

**Collections:** ATCC CRL 1751; ECACC 87022501; DSM (ACC 25); ICLC ATL 96004; SPBIC.



**Origin:** rat Wistar, sarcoma, subline of cell line XC derived from sarcoma, induces in vivo by Raus sarcoma

Submitted from Cardiological Scientific Centre. Moscow. 1979

**Morphology:** epithelial-like

**Mode of cultivation:** monolayer

**Conditions for cultivation:** medium - EMEM  
serum - FBS 10%

subculture procedure - cells detach from flask using trypsin 0.25%: EDTA 0.02% (1:3), split ratio 1:4 - 1:6

cryoconservation - growth medium, 10% DMSO,  $1.0 \times 10^6$  cells/ml in ampule

**Viability after cryoconservation:** 98% (0 passage, dye trypan blue)

**Sterility:** tests for bacteria, fungi and mycoplasma were negative

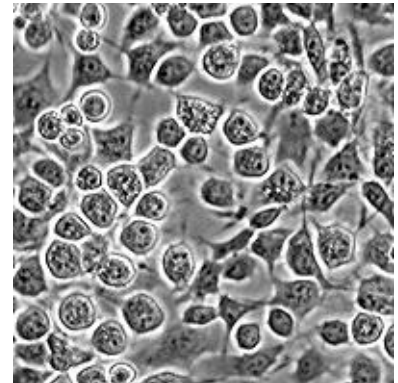
**Species:** karyological and immunofluorescent analysis

**Karyology:**  $2n=42$ , variability in the range between 40-45 chromosomes, modal number of chromosomes 42-43, number of markers -10 (differential dye), number of polyploid cells 70%

**Plating efficiency:** 68 %

**Applications:** cell biology

**Collections:** SPBIC.



**Origin:** mouse A/Sn, lymphoma induced in vivo by MLV.  
Eur. J. Immunol. 1975. 5: 112-117.

**Morphology:** lymphoblast-like

**Mode of cultivation:** suspension

**Conditions for cultivation:** medium - RPMI 1640

serum - FBS 10%

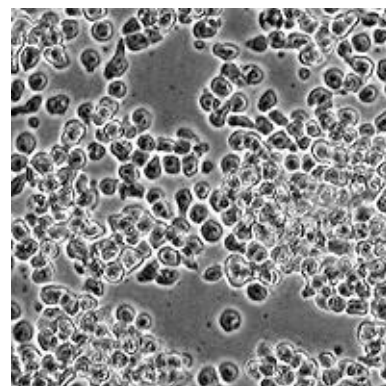
subculture procedure optimal

population density  $3.0-9.0 \times 10^5$  cells/ml

cryoconservation - growth medium,

10% DMSO,  $4.0-6.0 \times 10^6$  cells/ml in

ampule



**Viability after cryoconservation:** 80-90% (0 passage, dye trypan blue)

**Sterility:** tests for bacteria, fungi and mycoplasma were negative

**Species:** karyological and isoenzymological (LDH, G6PD) analysis

**Karyology:**  $2n=40$ , variability in the range between 40-47 chromosomes, modal number of chromosomes 43 without markers (routine and differential dye, C-banding), number of polyploid cells 2.5%.

**Other properties:** this cell line is sensitive to the cytotoxic activity of NK cells.

The cells not discovered of markers B- and T-lymphocytes (MWIIV).

**Applications:** NK assay, cytotoxicity.

**Collections:** ATCC TIB 160; ECACC 86022801; DSM ACC 96; MWIIV; SPBIC.