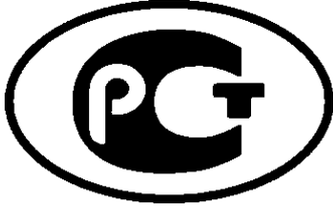


()

INTERSTATE COUNCIL FOR STANDARDIZATION, METROLOGY AND CERTIFICATION
(ISC)



ISO
14971 —
2011

(ISO 14971:2007, IDT)



2013

1.0—92 «
1.2—2009 «

1 «
- » ()

2

3

29 2011 . 40)

:

no MK (3166) 004—97	no (3166) 004—97	
	BY KZ KG RU	

4

2011 . 1261-

13

ISO 14971—2011

1 2013 .

5

Application of risk management to medical devices () .

ISO 14971:2007 Medical devices —

— (IDT).

14971—2009

6

« », — ()
« ».
« ».

1	1
2	1
3	3
3.1	3
3.2	5
3.3	5
3.4	5
3.5	6
4	6
4.1	6
4.2 , -	7
4.3	7
4.4 ()	7
5	8
6	8
6.1	8
6.2	8
6.3	9
6.4	9
6.5 /	9
6.6 ,	10
6.7	10
7	10
8	11
9	11
 ()	12
 ()	19
 () , -	21
 ,	25
D () ,	39
 ()	44
F ()	45
G ()	-
 ()	48
 <i>in vitro</i>	60
I () -	61
J ()	62
	62

ISO 14971:2007

/ 210 « -

/ 62 «
«
in vitro». / 212 «
in vitro»

a)

b)

J

I,

F,

in vitro.

Medical devices
Application of risk management to medical devices

— 2013—01—01

1

in vitro,

2

2.1

(accompanying document):

(1), 3.4.

2.2

(harm):

(2), 3.3).

2.3

(hazard):

(2), 3.5).

2.4

(hazardous situation):

(2), 3.6).

« ».

2.5

(intended use):

2.6
medical device):

in vitro (IVD) (*in vitro* diagnostic medical device) (IVD

2.7 (life-cycle): -

2.8 (manufacturer): -

1 « »

2 « » . [3], 3.6.

2.9 (medical device): -

in vitro -

- ;

- ;

- ;

- ;

- ;

- ;

- in vitro ,

- ;

- ;

- ;

1 (Global Harmonization

Task Force — GHTF) (. (4)).

2 ,

- :

- ;

- / (. 3);

- ;

- ;

3 , -

- -

2.10 (objective evidence): -

- .

- ,

- (. [5], 3.8.1).

2.11 (post-production): -

- .

- :

- , , , ,

- .

2.12 (procedure): -

(. [5], 3.4.5).

2.13 (process): ,

(. [5]. 3.4.1).

2.14 (record): -

(. [5], 3.7.6).

2.15 (residual risk): ,

1 (2). 3.9.
2 [2], 3.9, « » « ».

6.2.

2.16 (risk): (. [2], 3.2).

2.17 (risk analysis):
(. [2], 3.10).

2.18 (risk assessment): (. [2], 3.12).

2.19 (risk control):

2.20 (risk estimation): ,

2.21 (risk evaluation): , ,

2.22 (risk management): , , ,

2.23 (risk management file): ,

2.24 (safety): (. [2], 3.1).

2.25 (severity):

2.26 (top management): ,

— (5), 3.2.7.

2.27 (use error): ,

(. [6], 2.12).

1 (. [6], 2.12).

2 . [6J, D.1.3.

3

2.28 (verification):

(. [5], 3.8.4).

1 « »

2 :

- ;

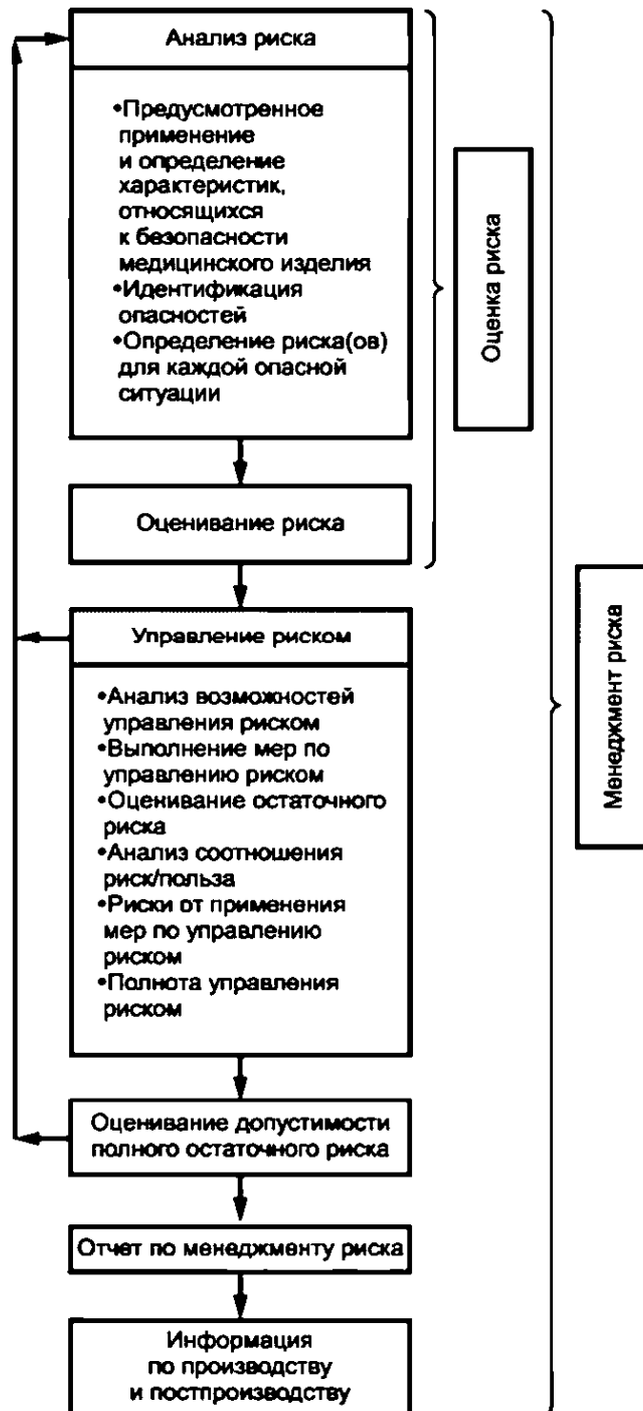
• ;

- ;

-

3

3.1



1

2

1.

-
-
8

3.2

(3.3)

3.3

()

3.4

)

, 8

b)

c)

d)

e)

f)

1

2

3

•

-

,

(3.2).

3.5

-

-

-

-

1

2

4

4.1

4.2—4.4.

1

2
3
in vitro
4

G.

I.

4.2—4.4.

a)
b)
c)
5

(

).

4.2

1

2

4.3

() .

.2 () .24

4.4

()

() .

() ()

1

(G).

2

.2.4.5 () .4 ().

3

()

(),

1

D)].

2

D.
in vitro.

3

a)

b)

c)

d)

e)

0

)

h)

5

6.2—6.6

6.7).

1

D.4 (D).

2

6.3 6.6.

6

6.1

6.2—6.7.

6.2

()

()

- a) ;
 - b) ;
 - c) .
- 1)), *
- 2 , , -
- 3 , , -
- 4 , (, -
- 5 D). , D.3.2.3
- J. -

/ (6.5). ,

6.3

6.2.

6.4

i () () () () , (. 6.2).

J. -

()

6.5 /

6.6.

— D.6 (D).

6.6

- a)
- b)

4.4—6.5.

6.7

()

7

(D).

0.7

J.

()

8

• ; ; :
- ;
- ;
8
[3.4,)].

9

a) ; / ,
b) ;
(), () ;
() () ;

1

2 [3]. 8.2.

()

» 1 / 210 — / 62 «

ib i i ,

4 / 210.

15 / 62 . 1 (JWG 1)

) (

« »

(— D)

.2

.2.1

in vitro

.2.2

[2], [3], [5].

1 (JWG 1)

« » (2.2)

« ,

« » (2.8),

[10],

(2.9)

[3],

(GHTF) [4].

» (2.5)

« » () « » ().

» (2.11), «

» (2.19). «

» (2.21), «

» (2.7), «

» (2.20). «

» (2.22) «

» (2.23).

« »

[2].

: 1)

; 2)

« » « » [7].

« » (2.26) [5].

.2.3
.2.3.1

3.1.

()

3.2 3.3.

()

3.2 3.3

.2.3.2

- a)
- b)
- c)
- d)

(.2.3.3);

.2.3.3

-
-
-
-
-

.2.3.4

- a)
- b)

)
)—f) (. 3.4)

6.3.

.2.3.5

« »

.2.4

.2.4.1

4.1

1 (. 4.1)

),)) (. 4.1, 4)

) (. 4.1, 4),

.2.4.2

()

, «
in vitro
in vitro»

/ 212 «

».

.2.4.3

4.2.

.2.4.4

()

() () (. .1 ())].

D

[8].

(8),

D

.2.5

5

.2.6

.2.6.1

6.2-6.7

.2.6.2

- a)
- b)
- c)

(2]

(9].

(, (10)).

2.6.2 /

6.5,

.2.6.3

—

.2.6.4

.2.6.5

6.5

(J).

.2.6.6
6.6

.2.6.7

.2.7

4—6,

.2.8

.2.9

, 3.5

1 (JWG 1).

» «

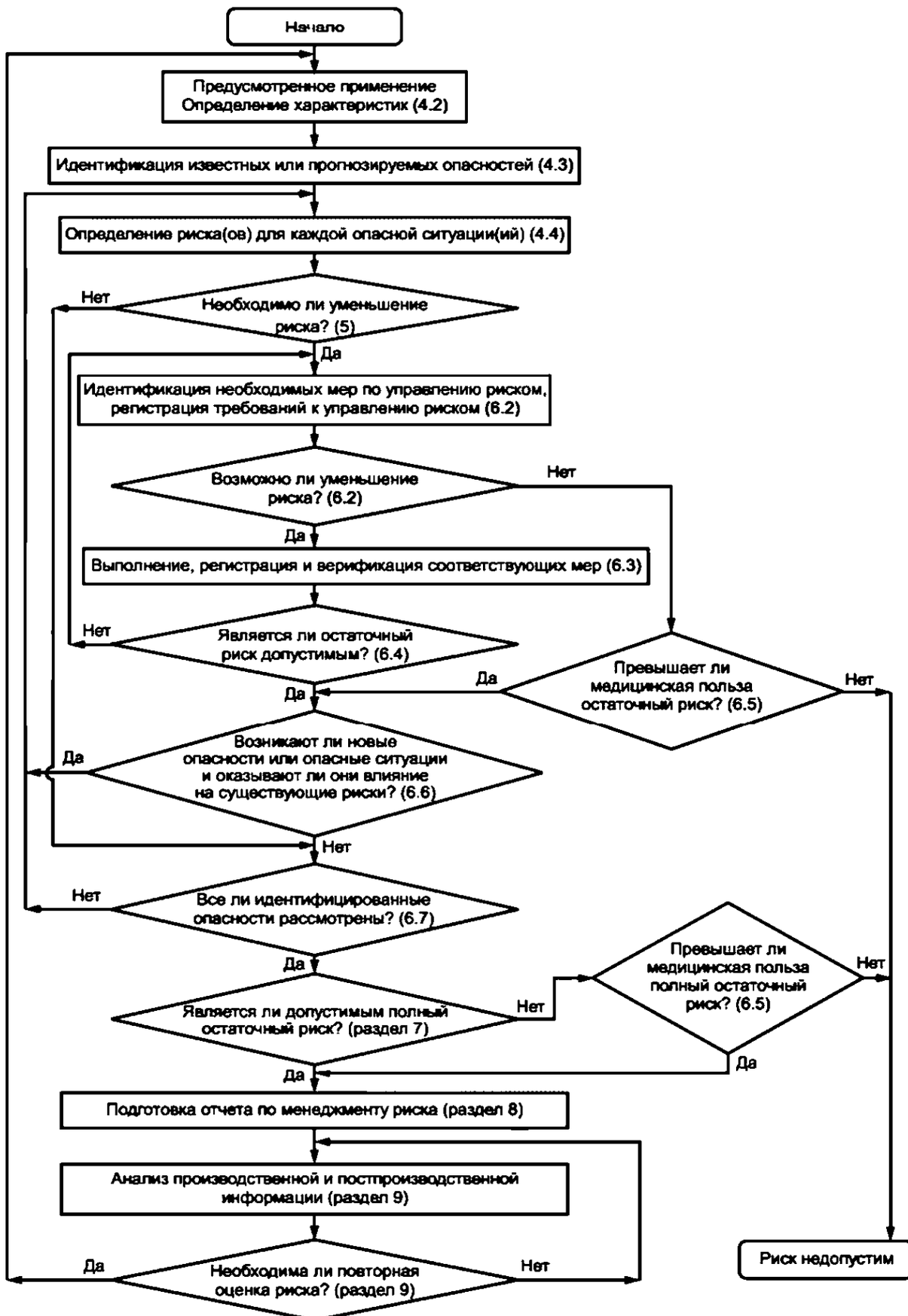
»

«

9

()

.1 , . .1. -
, , , -
.



()

.1

4.2.

(. 4.3).

in vitro, .2.5.4 ().

.2

.2.1

?

.2.2

?

.2.3

?

.2.4

?

(11).

.2.5

?

.2.6

?

.2.7

.2.8

.2.9

.2.10

.2.11

.2.12

.2.13

tdMH

1 ?

.2.14

.2.15

.2.16

.2.17

.2.18

.2.19 ?

.2.20 ?

.2.21 ?

.2.22 ?

.2.23 ?

.2.24 ?

.2.25 ?

.2.26 ?

.2.27 ?

.2.28 ?

.2.29 ?

.2.29.1 ?

.2.29.2 ?

.2.29.3 ?

[13].

[12].

(D)

D.1

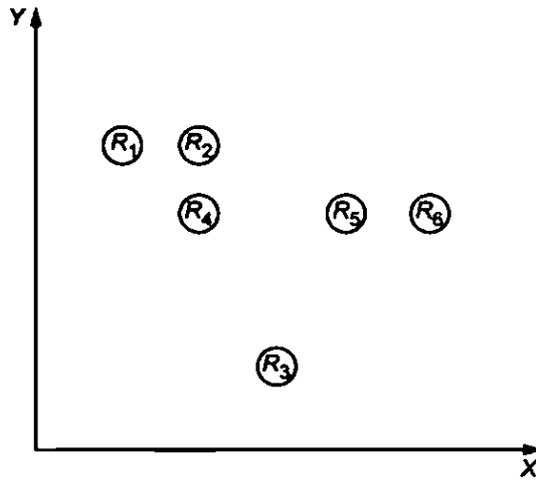
-
-
-
-
-
-

« » 2.16 « ».

X

D.1.
Y.

8
R₂.R₃. •)



* X—

; * Y—

D.1 —

D.2
D.2.1

in vitro.

D.2.2
D.2.2.1

D.2.2.2

D.2.2.3

D.3
D.3.1

in vitro)'

(. 0.2.2.3).

(14).

IVD- (

(. 4.4).

D.3 J.3

D.2.2.3,

D.3.2.3,

(. D.4).

TM

D.3.3

(S1. S2),

R
0.3.4).

D.3.4
D.3.4.1

NxM

N

3x3.

D.1 0.2.

« - » ,
 - -
 : -
 - , : ;
 -
 -
 , -
 ,
 : , D.4.
 D4
 0.5. ()

	*1	r ₂	
			*5-«6
		*3	

- — : ;- —
 D.4 — 3x3

	-	-		-	-
		r ₂			
		*4		*5	*6
			*3		

- — : ;- —
 0.5 —

(, « » .), (.0.8.5).
 D.5
 D.5.1

() /
 :

a)

-
-
-

b)

-
-

;

c)

-
-
-

-
-

-
-

-
-

-
-

-
-

-
-

-
-

-
-

-
-

-
-

-
-

-
-

-
-

-
-

-
-

-
-

-
-

-
-

-
-

-
-

-
-

-
-

-
-

-
-

-
-

-
-

-
-

-

D.5.2

D.5.3

D.5

D.5 —

/	-	-	,	/	
- - -		(-)		-	-
		-	- - -	- - -	
- - -	-	- - -	- -	- -	- - -

D.5

/	-	-	,	/	
-	-		-	-	-
-	-	(-	-	-
	,	-			
)			

D.5.4

Analysis on Critical Control Points — HACCP [c.m. G.6 (D.5.5) and G.7]. (Hazard)

D.6
D.6.1

(. D.8.4)

D.2

D.7.3

()

«

»

«

»,

D.7.4

). 8

(

D.7.5

8

D.7.6

D.7.7

(

).

D.7.8

8

. 8

D.8

D.8.1

«

»

» (ALARP-).

«

a)

b)

c)

));

» (ALARP-)

(. 6.2). ALARP-

BJ2

D.8.3

D.8.4

ALARP-

).

ALARP-

B.5

0.6

(Rj, /?2< ^ >)•

	-	-		-	-
		«2			
		*4		*5	
			*3		

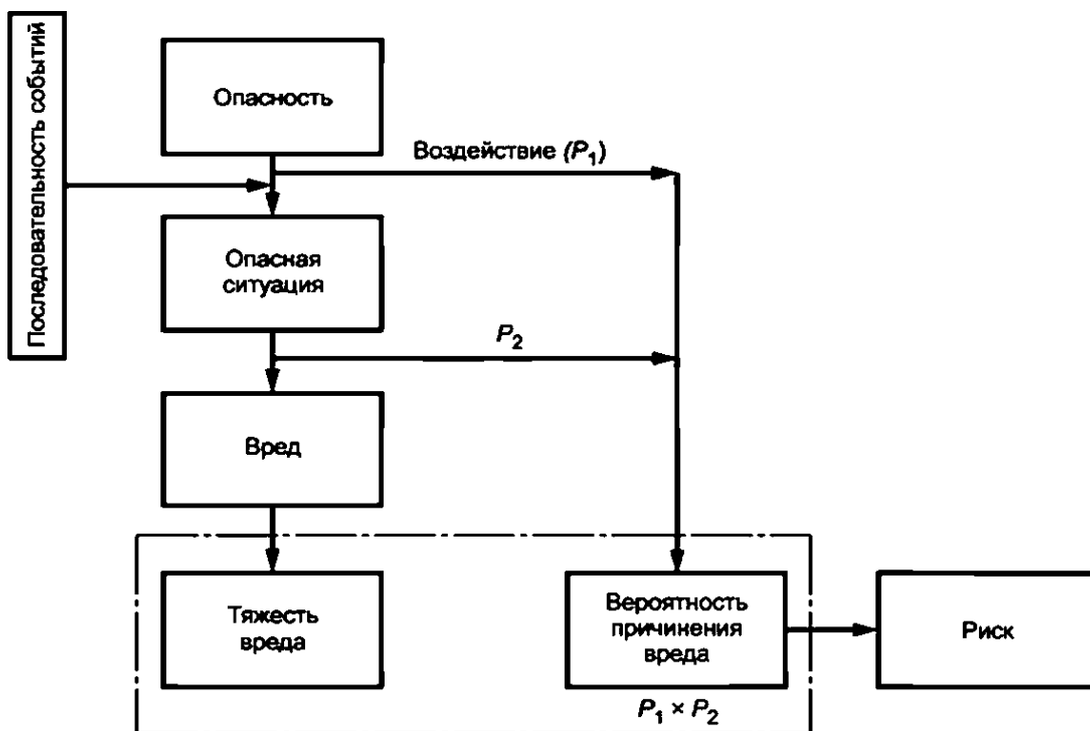
- - - - ; ;

D.6 —

.1

4.3.

4.4.



• 2 —

.1—

8

.1.

()			
()			
()			
(ESD —)	ESD		
()			

(F)

F.

8 3.4.

3.4.

F 2

(. [3]),

Б.

() ()

(), (),
(. 3.2).

В.

(. [3], 7.3.4).

Б.

[. D.4 (D)].

[3]. 7.1).

Б.

[. .2.6.3 ()].

2.26).

F. ()
()

(. [3]. 8.2).

(G)

G.

4.3.

(Preliminary Hazard Analysis — PHA) —

(Fault Tree Analysis — FTA) —

(Failure Mode and Effect Analysis — FMEA)
(Failure Mode, Effects and Criticality Analysis — FMECA)

(Hazard and Operability Study — HAZOP)
(Hazard Analysis on Critical Control Points — HACCP) —

B.

()
() —

- a)
- b)
- c)
- d)
- e)

[8]. 5.

G.

FTA

(FTA)

FTA

FTA [17].
(FMEA)
(FMEA)

6.

: « ...?».

« », . . .

FMEA

FMEA). FMEA

(FMEA)

FMEA

(FMECA).

FMECA

FMEA

[18].
(HAZOP)
(HAZOP)

6.

FMEA.

HAZOP

),

a)

b)

. .)

(NONE — ; PART OF —

HAZOP [19].
()
()—

6.

(NASA)

()

in vitro

.1

in vitro (IVD).

in vitro.

in vitro

in vitro,

in vitro

IVD-

IVD-

in vitro

ib

in vitro,

« i » «

().

« »,

in vitro-

in vitro

in vitro,

in vitro,

.1

in vitro

()

()

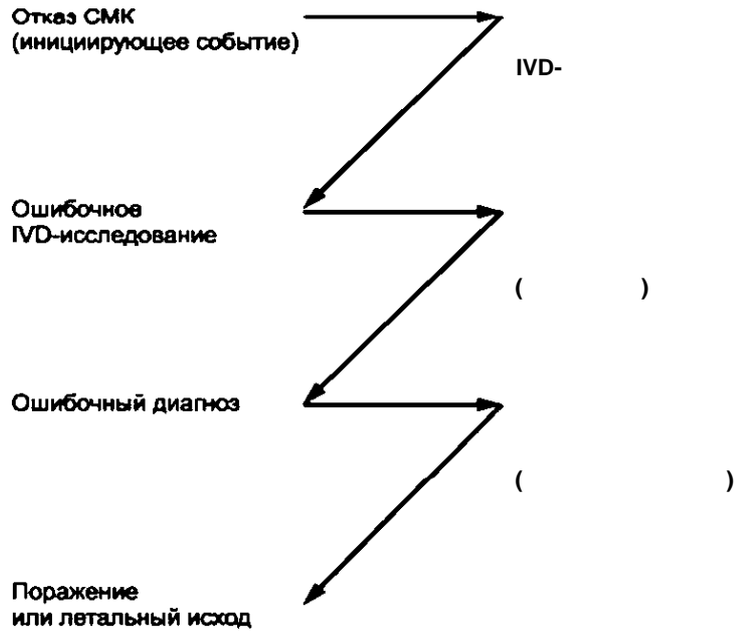
in vitro.

IVD-

in vitro-

.1.

in vitro



.1 —
in vitro

.1

.2
.2.1
.2.1.1

in vitro,

IVD-

: 1)

IVD-

; 2)

in vitro-

«

»

IVD-

.2.1.2

in vitro

(P-hCG)

Ji-hCG

.2.1.3

in vitro.
p-hCG

.2.2

.2.2.1

.2.2.2

in vitro

.2.2.3

in vitro-

IVD-

IVD-

IVD-

IVD-

IVD-

.2.2.4

.2.3

.2.3.1

in vitro

.2.3.2

.2.3.3

.2.3.4

in vitro

.2.3.5

in vitro
(. .)

.2.4

.2.4.1

IVD-

IVD-

in vitro,

.1.

(,)

(,),

(. . .2.3.2 .2.3.3);

(. . .2.3.4);

(. . .2.3.5).

.2.4.2

(. . .2.3),

(Preliminary Hazard Analysis

(Fault Tree Analysis — FTA).

(Failure Mode and Effect Analysis — FMEA)

(Hazard Analysis on Critical Control Points — HACCP),

6.

H.2.4.3

IVD-

(, , , . .). :

... « »), (-
: 1) ; 2) -
; 3) () , -
).

.2.5.2

IVD-

.2.1 .2.2.

IVD-

IVD-

.2.5.3

in vitro

in vitro

.1,

in vitro

() ;

in vitro

in vitro

.2.5.4.

in vitro,

in vitro.

.2.5.4

.2.5.4.1

in vitro:

.2.5.4.2

IVD-

in vitro;

);

» ;

.2.5.4.3 IVD- ;

;

;

);

;

.2.5.4.4 ()

in vitro

in vitro

in vitro

.2.5.4.5 ;

;

.2.5.4.6 :

;

;

();

.2.5.5 *in vitro*

.2.5.5.1

IVD- *in vitro*,

.2.5.5.2

[21]

[22].

.2.5.5.3

[21],

IVD-

; 2)

: 1)

; 3)

; 4)

D.3 D.4 (D).

.4
.4.1

).

IVD-
IVD-

IVD-

6.2.

in vitro

a)

(

b)

c)

1

2

in vitro,

(. 4.2.4).

.4.2
.4.2.1

in vitro

- ;
 - ;
 - IVD- (, ;);
 - (,);
 • ;
 - ;
 - (, -);
 - (,).

in vitro,
).

(. . -
[Hazard Analysis

on Critical Control Points — HACCP, c.m. G.6 (G)J

, :
 - ;
 - , ;
 ;
 - ,

4.2.2

in vitro

8

. , : ,
 - (,) ;
 -) : (,) ;
 - (, , ,) ;
 • , ;
 ;

(. [23] [24]);

4.2.3

4.2.3.1

in vitro,

. .
 : (,) ;
 - ; (. . (,)) ;
 - ;
 - (()) ;
 - () ;
 - () .

4.2.3.2

... () ;
 ;
 ;
 ;
 ;
 ;

4.2.3.3

[25];
 () ;
 HPLC- GC- () ;
) ;

4.2.3.4

in vitro
in vitro () ;
 (. [26]).

4 J.4

MHoiMx ci * 1
in vitro.

4.3).

.1

.1 —

	(, , . .)

in vitro

IVD-

.5
.5.1

in vitro

in vitro

(External Quality Assessment Schemes

— EQAS),

.5.2

in vitro

[1]	IEC 60601-1:2005	Medical electrical equipment — Part 1: General requirements for basic safety and essential performance (. 1.)	-
[2]	(ISO/IEC Guide 51:1999)	Safety aspects — Guidelines for their inclusion in standards (. 8)	-
[3]	ISO 13485:2003	Medical devices — Quality management systems — Requirements for regulatory purposes (.)	-
[4]	(Document No. N029R11, dated 2 Feb. 2002)	Global Harmonization Task Force (GHTF) — Study Group 1 (SG1) (no 1)	-
[5]	ISO 9000:2005	Quality management systems — Fundamentals and vocabulary (.)	-
[6]	IEC 62366:2007	Medical devices - Application of usability engineering to medical devices (.)	-
[7]	IEC 60601-1-4:1996	Medical electrical equipment — Part 1-4: General requirements for safety — Collateral standard: Programmable electrical medical systems (. 1-4.)	-
[8]	IEC 60300-3-9:1995	Dependability management — Part 3: Application guide — Section 9: Risk analysis of technological systems (. 3. 9.)	-
[9]	IEC/TR 60513:1994	Fundamental aspects of safety standards for medical electrical equipment (.)	-
[10]	(Council Directive 93/42/E EC of 14 June 1993)	Directive concerning medical devices as amended by Directive 98/79/EC of the European Parliament and of the Council of 27 October 1998 on <i>in vitro</i> diagnostic medical devices (. (. , 98/79/ <i>in vitro</i>))	-
[11]	ISO 22442:2007 (all parts)	Medical devices utilizing animal tissues and their derivatives (.)	-
[12]	IEC 60601-1-6:2010	Medical electrical equipment — Part 1-6: General requirements for basic safety and essential performance — Collateral standard: Usability (. 1-6.)	-
[13]	IEC 60601-1-8:2006	Medical electrical equipment — Part 1-8: General requirements for basic safety and essential performance — Collateral standard; General requirements, tests and guidance for alarm systems in medical electrical equipment and medical electrical systems (. 1-8.)	-
[14]	ISO 10993-17:2002	Biological evaluation of medical devices — Part 17: Establishment of allowable limits for leachable substances (. 17.)	-
[15]	ISO 14155-1:2003	Clinical investigation of medical devices for human subjects — Part 1; General requirements (. 1.)	-

[16] ISO 14155-2:2003 Clinical investigation of medical devices for human subjects — Part 2: Clinical investigation plans
() 2. -

[17] IEC 61025:2006 Fault tree analysis (FTA)
()

[18] IEC 60812:2006 Analysis techniques for system reliability — Procedures for failure mode and effects analysis (FMEA)
() -

[19] IEC 61882:2001 Hazard and operability studies (HAZOP studies) — Application guide (HAZOP).
() -

[20] ISO 15197:2003 *In vitro* diagnostic test systems — Requirements for blood-glucose monitoring systems for self-testing in managing diabetes meNitus
(*in vitro*.) -

[21] (Parkes J.L. et al., Diabetes Care. 23. 2000) A new consensus error grid to evaluate the clinical significance of inaccuracies in the measurement of blood glucose
() -

[22] (Clarke W.L. et al., Diabetes Care. 10(5), 1987) Evaluating Clinical Accuracy of Systems for Self-Monitoring of Blood Glucose
()

[23] ISO 17511:2003 *In vitro* diagnostic medical devices — Measurement of quantities in biological samples — Metrological traceability of values assigned to calibrators and control materials
(*in vitro*.) -

[24] ISO 18153:2003 *In vitro* diagnostic medical devices — Measurement of quantities in biological samples — Metrological traceability of values for catalytic concentration of enzymes assigned to calibrators and control materials
(*in vitro*.) -

[25] ISO 15198:2004 Clinical laboratory medicine — *In vitro* diagnostic medical devices — Validation of user quality control procedures by the manufacturer
(*in vitro*.) -

[26] ISO 17593:2007 Clinical laboratory testing and *in vitro* diagnostic test systems — Requirements for *in vitro* monitoring systems for self-testing of oral-anticoagulant therapy
(*in vitro*) *in vitro*. -

[27] ISO 19001:2002 *In vitro* diagnostic medical devices. Information supplied by the manufacturer with *in vitro* diagnostic reagents for staining in biology
(*in vitro*.) *in vitro* -

[28] ISO 10993-1:2009 Biological evaluation of medical devices — Part 1: Evaluation and testing within a risk management system
() 1. -

617.7-08.001.33:006.354

11.040.01

IDT

: , ,

• •
• •
• •
• •

02.09.2013.

24.10.2013.

60x84%.

. . 7.91. . 6.47. 51 . 1286

« », 123995 . ., 4.

www.gostinfo.njinfo@gostinfo.ru

. 248021 , . . 256.