

Timeframes' monitoring (2014 year)

The survey was carried out in January-February 2015.

The sample included submissions made from January 1, 2014 till December 31, 2014 as well as submissions made before January 1, 2014 if decisions on granting/refusing approvals were taken in 2014.

24 ACTO member companies took part in the survey (pharmaceutical companies and contract research organizations)...

Table 1 (Timeframes for Issuing Approvals) provides information on average, minimum and maximum timeframes for issuing approvals to conduct clinical trials, permits to import medicines for clinical trials, permits to import/export biological materials, approvals to make amendments to the protocols and other approvals (to prolong clinical trials, to involve new sites, to enroll additional patients, etc.)

The total time to begin a trial is the sum of average timeframes for issuing approvals to conduct clinical trials and average timeframes for issuing permits to import/export biological materials.

Actual timeframes are given in calendar days. Timeframes according to legislation are given in calendar as well as in business days.

Table 2 (Changes in average timeframes for issuing approvals to conduct clinical trials and permits to import/export) provides information on average timeframes for issuing approvals to conduct clinical trials and permits to import/export as well as total time to begin a trial in 2005-2014. It's worth noting that before 2011 year ACTO didn't differentiate timeframes for issuing permits to import/export biological materials and medicines.

Before September 2010 examinations carried out by FGU and Ethics Committee weren't part of the approval process. That's why average timeframes for issuing approvals to conduct clinical trials in 2004-2010 were calculated as the sum of average timeframes for carrying out examinations by Ethics Committee and FGU (the highest number was taken into account) and timeframes for issuing approvals by Roszdravnadzor.

Table 3 provides information on violations of timeframes.

Tables 4-8 provide detailed information on timeframes for issuing all sorts of approvals. Actual timeframes are given in calendar days. Timeframes according to legislation are given in business and calendar days. The column «Legislation» provides information on relevant sources of the numbers in the column «Timeframes according to legislation».

The column «Sampling» provides information on the number of submitted applications. The samples are different at different stages because companies apply various approaches to timeframes monitoring.

Table 1. Timeframes for Issuing Approvals

Table 1. Tilliellallies for issuing A	ppiovaio				
	Timeframes according to legislation (business/cal endar days)	Average timeframes (calendar days)	Minimum timeframes (calendar days)	Maximum timeframes (calendar days)	Sampling
To conduct clinical trials	41/57	95	30	244	165
To import medicines	8/12	14	3	48	350
To import/export biological materials	13/19	23	2	62	645
To make amendments to the protocol	34/48	60	7	105	324
Other approvals (to prolong clinical trials, to involve new sites, to enroll					
additional patients, etc.)	25/35	27	6	109	634
Total time to obtain approvals to conduct clinical trials and to import/export*	54/76	118	~	2	~

^{*} It's the sum of average timeframes for issuing approvals to conduct clinical trials and average timeframes for issuing permits to import/export biological materials or to import drugs (the highest number was taken into account)

Table 2. Changes in average timeframes

	2005	2006	2007	2008	2009	Jan-Aug 2010	2011	2012	2013	2014
Approvals to conduct clinical										
trials	66,3	77,8	98,9	77,6	77,0	85,2	130,0	116,0	87,0	95,0
Permits to import/export	14,9	17,8	23,7	33,1	30,5	26,9	34,0	20,0	20,0	23,0
Total	81,2	95,6	122,6	110,7	107,5	112,1	164,0	135,0	107,0	118,0

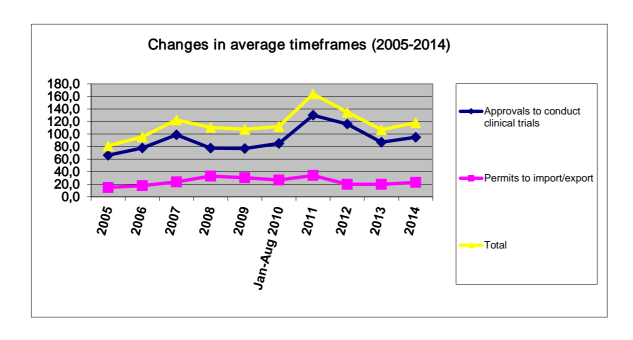


Table 3. Violations of timeframes

Table 3. Violations of timeral			Appro	ovals issued in v	violation of time	eframes	
	Approvals issued on time	Total	less than in 1,5 times	in 1,5-1,9 times	in 2-2,9 times	in 3-3,9 times	In 4 times and more
to conduct clinical trials	6,1%	93,9%	44,9%	43,0%	4,8%	1,2%	0,0%
to import medicines	42,0%	58,0%	33,4%	16,6%	7,1%	0,6%	0,3%
to import/export biologocal materials	29,5%	70,5%	43,5%	24,5%	2,3%	0,2%	0,0%
to make amendments to the protocol	25,0%	75,0%	44,7%	28,4%	1,9%	0,0%	0,0%
Other Approvals (to Prolong Clinical Trials, to Include New Sites, to Enroll Additional Patients, etc.)	86,8%	13,2%	10,6%	1,9%	0,5%	0,2%	0,0%

Table 4. Timeframes for Issuing Approvals to Conduct Clinical Trials

Table 4. Timetrames for Issuing Approvals to	Timeframes according to				a				
Stages of review	legislation (business/calendar days)	Legislation (references)	average timeframes	minimum timeframes	1st quartile	median	3rd quartile	maximum timeframes	Sampling*
Application registration (from the date of the submission)	1/1	Paragraph 16 of the Rules of proceedings in the federal executive bodies (Government Order of June 15, 2009 N 477)	1	0	0	0	1	26	210
Data completeness check and decision on issuance of an assignment to carry out expert examination (from the date of the registration)	5/7	Peragraph 3 of the Article 39 of the Federal Law On Circulation of Medicines	5	0	2	4	6	31	179
Examinations (from the date of the decision)	30/42	Paragraph 1 of the Article 20 of the Federal Law On Circulation of Medicines	55	20	49	54	61	89	64
Second application submission (from the date of the first application submission)	~	There is no such requirement in the legislation	80	29	57	68	84	292	150
Decision on issuance of an approval to conduct a clinical trial	5/7	Paragraph 2 of the Article 22 of the Federal Law On Circulation of Medicines	8	1	5	7	8	33	149
Receipt of the decision on issuance of an approval to conduct a clinical trial (from the date of the decision on issuance of an approval)	0/0	Paragraph 2 of the Article 22 of the Federal Law On Circulation of Medicines	12	3	8	11	14	38	168
Total time to receive a notification that a permit is refused (in case of refusal) from the date of the submission	41/57	the sum of all stages	71	28	58	69	75	196	47
Total time to receive a permit from the date of the submission	41/57	the sum of all stages	92	28	61	74	111	213	47
Total time to receive a permit from the date of the resubmission (if initial submission was refused)	41/57	the sum of all stages	84	30	72	82	93	177	165
Total time to receive a permit after resubmission from the date of the initial submission (if initial submission was refused)	41/57	the sum of all stages	95	30	75	87	104	244	165

^{*}The samples are different as different companies apply different approaches to timeframes' monitoring

Table 5. Timeframes for Issuing Permits to Import Medicines

Stages of review	Timeframes according to legislation Legislation (references)		Practice (calendar days) total number of applications is 353						
	(business/calendar days)	Legislation (references)	average timeframes	minimum timeframes	1st quartile	median	3rd quartile	maximum timeframes	Sampling*
Application registration (from the date of the submission)	1/1	Paragraph 16 of the Rules of proceedings in the federal executive bodies (Government Order of June 15, 2009 N 477)	0	0	0	0	0	6	128
Data completeness check and decision on whether to grant a permit or to notify that a permit is refused (from the date of the registration)	5/7	Paragraph 12 of the Rules of Import of Medicines for Medical Use into the Russian Federation (Government Order of September 29, 2010 № 771)	7	0	4	6	9	25	129
Receipt of the decision (from the date of the decision)	2/4	Paragraph 22 of the Rules of proceedings in the federal executive bodies (Government Order of June 15, 2009 N 477)	8	0	5	7	10	33	339
Total time to receive a permit or a notification that a permit is refused (from the date of the submission)	8/12	the sum of all stages	14	3	11	13	17	48	350

Table 6. Timeframes for Issuing Permits to Import/Export Biological Materials

Stages of review	Timeframes according to legislation	Legislation (references)	Practice (calendar days) total number of applications is 646						
	(business/calendar days)	Degisiation (references)	average timeframes	minimum timeframes	1st quartile	median	3rd quartile	maximum timeframes	Sampling*
Application registration (from the date of the submission)	1/1	Paragraph 16 of the Rules of proceedings in the federal executive bodies (Government Order of June 15, 2009 N 477)	0	0	0	0	0	7	325
Data completeness check and decision on whether to grant a permit or to notify that a permit is refused (from the date of the registration)	10/14	Paragraph 5 of the Rules for Import and Export of Biological Materials into and outside of the Russian Federation (Government Order of September 3, 2010 Ne673)	13	0	11	13	14	38	329
Receipt of the decision (from the date of the decision)	2/4	Paragraph 22 of the Rules of proceedings in the federal executive bodies (Government Order of June 15, 2009 N 477)	9	0	6	8	12	42	614
Total time to receive a permit or a notification that a permit is refused (from the date of the submission)	13/19	the sum of all stages	23	2	18	22	28	62	645

^{*}The samples are different as different companies apply different approaches to timeframes' monitoring

Table 7. Timeframes for Issuing Approvals to Make Amendments to the Protocol

Stages of review	Timeframes according to legislation	I arialetian (arfananasa)	Practice (calendar days) total number of applications is 333						
	(business/calendar days)	Legislation (references)	average timeframes	minimum timeframes	1st quartile	median	3rd quartile	maximum timeframes	Sampling*
Application registration (from the date of the submission)	1/1	"Paragraph 16 of the Rules of proceedings in the federal executive bodies (Government Order of June 15, 2009 N 477)	0	0	0	0	0	11	127
Decision on whether to grant an approval or to notify that an approval is refused (from the date of the registration)	30/42	Paragraph 5 of the Article 40 of the Federal Law On Circulation of Medicines, Paragraph 7 of the MoH Order of August 31, 2010 N 775n	52	4	44	55	63	92	149
Receipt of the decision (from the date of the decision)	3/5	Paragraph 10 of the MoH Order of August 31, 2010 N 775n	9	0	6	7	11	38	302
Total time to receive an approval or a notification that an approval is refused (from the date of the submission)	34/48	the sum of all stages	60	7	51	62	72	105	324

Table 8. Timeframes for Issuing Other Approvals (to Involve New Sites, To Enroll Additional Patients, to Prolong Clinical Trials etc.)

Stages of review	Timeframes according to legislation	Legislation (references)	Practice (calendar days) total number of applications is 636						
Stages of Feview	(business/calendar days)	Legislation (references)	average timeframes	minimum timeframes	1st quartile	median	3rd quartile	maximum timeframes	Sampling*
Application registration (from the date of the submission)	1/1	Paragraph 16 of the Rules of proceedings in the federal executive bodies (Government Order of June 15, 2009 N 477	1	0	0	0	1	11	297
Decision on whether to grant an approval or to notify that an approval is refused (from the date of the registration)	22/30	Article 12 of the Federal Law On Russian citizens' requests consideration of May 2, 2006 № 59	19	3	13	18	23	94	297
Receipt of the decision (from the date of the decision)	2/4	Paragraph 22 of the Rules of proceedings in the federal executive bodies (Government Order of June 15, 2009 N 477)	8	0	5	7	9	46	597
Total time to receive an approval or a notification that an approval is refused (from the date of the submission)	25/35	the sum of all stages	27	6	21	26	32	109	634

^{*}The samples are different as different companies use different approaches to timeframes' monitoring