Summaryof the working program of the academic discipline

«Toxicological chemistry» (name of the academic discipline) General Educational Program of higher education (<u>specialist's degree programs</u>) 33.05.01 "Pharmacy" Department: Pharmaceutical Chemistry and Pharmacognosy

1. The purpose of mastering the discipline: participation forming the relevant competencies (UC-1, UC-2), general professional (GPC-1, GPC -3, GPC -6) and professional (PC-4, PC-7, PC-12) competencies

2. Position of the academic discipline in the structure of the General Educational Program (GEP).

2.1. The discipline Toxicological chemistry refers to the core part (or the part formed by the participants of educational relations) of Block 1 of GEP HE (Academic discipline index). The discipline is taught in 7, 8 semesters.

3. Deliverables of mastering the academic discipline and metrics of competence acquisition

Mastering the discipline aims at acquiring the following universal (UC) or/and general professional (GPC) or/and professional (PC) competencies

	Competence	The content of	Code and name of	As a result of ma	astering the discipli should:	ne, the students
№	Competence code		the competence acquisition metric	know	be able to	possess
1.	UC-1.	Able to realize critical analysis of problem situations based on a systematic approach, develop strategy actions	UC-1.1. Analyzes the problem situation as a system identifying its components and connections between themUC- 1.2. Identifies gaps in the information needed to solve a problem situation, and designs processes for their elimination UC-1.3. Critically assesses reliability of information sources, works with conflicting information from different sources UC-1.4. Develops and meaningfully argues the strategy of solving the problem situations based on the system and interdisciplinary approaches UC-1.5. Uses logical and	 methodology of abstract thinking for systematization of processes and construction of cause-and-effect relationships; modern theoretical and experimental methods for the implementation of own and borrowed results of scientific research into practice. 	 abstract, analyze and synthesize the information received; highlight and to systematize the essential properties and connections of objects, to identify the main patterns of the objects under study; search, select and analyze information obtained from various sources in order to make the best decision at the modern scientific level, in accordance with professional tasks and the requirements of legal documents. 	 methods of self-control, abstract and analytical thinking; skills in analyzing methodological problems that arise in solving research and practical problems, including those in interdisciplinary areas; skills of presenting an independent point of view

			methodological			
			tools for critical evaluation of modern concepts of philosophical and social nature in its subject areas			
2.	UC-2.	Able to manage the project at all stages of its life cycle	UC-2.1. Formulates a project task on the basis of the set problems and a method of its solutions through the implementation of the project management UC-2.2. Develops a project concept within the framework of the designated problem: formulates the purpose, tasks, justifies the relevance, significance, expected results and possible areas of their application UC-2.3. Plans necessary resources, including taking into account their replaceability UC-2.4. Develops a project implementation plan using planning tools UC-2.5. Monitors the progress of the project, corrects deviations, makes additional changes to the project implementation plan, clarifies zones of responsibilities of project participants	principles for developing a project implementation plan in the field of professional activity at all stages of its life cycle	develop a project implementation plan in the field of professional activity at all stages of its life cycle, providing for problem situations and risks	methods of planning and executing projects under conditions of uncertainty, managing the project (supporting the implementation of the project)
3.	GPC-1.	Able to use basic biological, physical- chemical, chemical, mathematical methods for	GPC-1.1. Applies basic biological methods of analysis for the development, research and examination of pharmaceuticals and medicinal plant raw	 organization of a system of state control over the production and manufacture of drugs; the main regulatory 	• apply chemical, biological, physico- chemical and other methods of analysis during the	 ensuring the process of quality control of medicines with equipment and consumables; basic chemical,

		the	materials	documents,	examination of	biological,
		development,	GPC-1.2. Applies	production and	medicines.	physico-
		research and	basic physical-	manufacture,		chemical and
		examination of medicines, the	chemical and	quality control,		other methods of
		manufacture of	chemical analysis methods for the	storage and use of medicines		analysis during the examination
		medicinal	development,	(domestic and		of medicines.
		products	research and	international		
			examination of	standards (GMP,		
			medicinal products	GLP, GCP, GPP),		
			and medicinal plant raw materials	pharmacopoeias, orders of the		
			GPC-1.3. Applies	Ministry of		
			the basic methods of	Health of the		
			physical-chemical	Russian		
			analysis in the	Federation,		
			manufacture of	guidelines and		
			medicinal products GPC-1.4. Applies	instructions approved by the		
			mathematical	Ministry of		
			methods and	Health of the		
			performs	Russian		
			mathematical	Federation) for		
			processing of data	examination using chemical,		
			obtained during the development of	biological,		
			medicines, as well	physicochemical		
			as research and	and other		
			examination of	methods;		
			medicines and	• pharmacopoeial methods of		
			medicinal plant raw materials	analysis used in		
			materials	the analysis of		
				medicinal		
				products using		
				chemical,		
				biological, physicochemical		
				and other		
				methods.		
4.	GPC-3.	Able to carry	GPC-3.1. Complies	• laws and	• put into	• skills in
		out professional	with norms and	legislative acts of	practice the	organizing and conducting
		activities	rules established by the authorized state	the Russian Federation,	basic principles of the system of	quality control
		taking into	authorities when	normative and	quality control	of medicines at
		account	solving the tasks of	methodological	and safety of	the level of their
		specific	professional activity	materials of the	medicines in the	production,
		economic, environmental,	in the field of medicine circulation	Ministry of Health of Russia,	conditions of	transportation and storage;
		social factors	GPC-3.3. Performs	regulating the	pharmaceutical organizations;	• the main
		within the	labor actions taking	procedure for	• to organize	methods of
		framework of	into account their	conducting	and carry out	pharmaceutical
		the system of	impact on the	examinations	the procedure	analysis
		regulations of the medicine	environment,	provided for in	for quality	provided for in the state
		circulation	preventing the occurrence of	the state registration of	control of medicines at the	the state registration of
		sphere	environmental	medicines;	level of their	medicines;
		-	hazards	•general	production,	• skills in
				principles of	transportation	carrying out
				development,	and storage	preventive
				testing and registration of	using methods of	measures to ensure the
				medicines;	pharmacopoeial	quality of
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				 the basic principles, strategies, methods and procedures for quality control of medicines in the conditions of pharmaceutical organizations used in the course of examinations provided for in the state registration of medicines, in accordance with the requirements 	analysis.	medicines at the level of their production, transportation and storage.
				of the current regulatory and legislative framework.		
5.	GPC-6.	Able to understand the principles of modern information technologies and use them to solve the tasks of professional activity	GPC-6.2. Performs an effective search for information necessary to solve the tasks of professional activity using legal reference systems and professional pharmaceutical databases GPC-6.3. Uses specialized software for mathematical processing of observational and experimental data in solving problems of professional activity	modern means of computing technology	use modern computer technology and basic office applications And graphic packages; evaluate way of implementing information systems and devices for solving task	methods of practical use modern to computers to search information processing and fundamentals numerical methods for solving applied tasks
6.	PC-4.	Able to participate in monitoring the quality, effectiveness and safety of medicines and medicinal plant raw materials	PC-4.1. Conducts pharmaceutical analysis of pharmaceutical substances, excipients and medicines for medical use of factory production in accordance with quality standards PC-4.2. Performs intra-pharmacy quality control of medical use manufactured in a pharmacy organization PC-4.3. Conducts pharmacognostic	 laws and legislative acts of the Russian Federation, regulatory and methodological materials of the Ministry of Health of Russia, regulating the procedure for quality control of medicines in the conditions of pharmaceutical organizations; methods of analysis used in the quality control of drugs in the conditions 	 apply chemical, physico- chemical methods of intra-pharmacy quality of drugs in the conditions of pharmaceutical organizations; draw up documentation of the established form for the control of manufactured medicinal products in the conditions of pharmaceutical 	 basic chemical and physico- chemical methods of intra- pharmacy quality control of drugs in the conditions of pharmaceutical organizations; registration of documentation of the established sample for the control of manufactured drugs in the conditions of pharmaceutical organizations.

			analysis of medicinal plant raw materials and medicinal herbal preparations PC-4.4. Informs in accordance with the procedure established by law about the non- compliance of the medicinal product for medical use with the established requirements or about the non- compliance of the data on the effectiveness and safety of the medicinal product with the data on the medicinal product with the data on the medicinal product with the data on the medicinal product contained in the instructions for its use	of pharmaceutical organizations; • monitor drug quality assurance systems; • the process of providing equipment and consumables for quality control in the conditions of pharmaceutical organizations;	organizations; • monitor drug quality assurance systems; • provide the process of quality control in pharmaceutical organizations with equipment and consumables.	
7.	PC-7.	Able to carry out operations related to the technological process in the production of medicines and their control	PC-7.5. Monitors the compliance of the raw materials and excipients used with the requirements of regulatory documentation	requirements of regulatory documentation for the raw materials and auxiliary materials used	carry out pharmacopoeial analysis of raw materials and auxiliary materials used	methods of quality control of raw materials and auxiliary materials used
8.	PC-12.	Able to take part in conducting chemical- toxicological and forensic- chemical research in order to diagnose poisoning, drug and alcohol intoxication	PC-12.1. Participates in conducting chemical and toxicological research PC-12.2. Conducts forensic chemical studies in order to detect poisoning, drug and alcohol intoxication PC-12.3. Participates in monitoring the quality and safety of medicines and medicinal plant raw materials	regulatory and legislative acts regulating the examination of poisoning and intoxication; • physico- chemical methods underlying the qualitative and quantitative analysis of poisonous substances in accordance with the requirements of the State Pharmacopoeia.	 apply the regulatory framework governing the examination of poisoning and intoxication; analyze poisonous substances, poisoning products, biological material in accordance with the requirements of regulatory documentation 	 apply the regulatory framework governing the examination of poisoning and intoxication; analyze poisonous substances, poisoning products, biological material in accordance with the requirements of regulatory documentation • skills in analyzing poisonous substances, poisoning products, biological material in accordance with the requirements

4. Volume of the academic discipline and types of academic work Total labor intensity of the discipline is 6 CU (216 AH)

Type of educational work	Labo	Labor intensity (AH) in		
	volume in credit	volume in credit units	semesters	
	units (CU)	(CU)		
			7	8
classroom work, including	3	108	66	42
Lectures (L)	0.7	24	14	10
Practicals (P)	2.3	84	52	32
Student's individual work (SIW)	2	72	42	30
Mid-term assessment				
exam	1	36		36
TOTAL LABOR CAPACITY	6	216	108	108

5. Sections of the academic discipline and competencies that are formed

№	Competence code	Section name of the discipline
1.	UC-1,2 GPC-1,3,6	Toxicological chemistry as a special discipline. Legal basis of chemical- toxicological analysis. Detoxification methods for acute poisoning
	PC-4,7,12	
	UC-1,2	Biochemical toxicology
2.	GPC-1,3,6	
	PC-4,7,12	
	UC-1,2	Analytical toxicology
3.	GPC-1,3,6	
	PC-4,7,12	