## FDA – Food and Drug Administration

Your Objectives:				
At the end of the lesson, you will be able to starrelation to the pharmaceutical industry.	te the basic role and importance of the FDA in			
The Food and Drug Administration (FDA)'s pri	ncipal mandate under the US Department of			
Health and Human Services is to protect public	of the USA. It			
administrates the	and efficacy of medicinal products used in			
both human and	medicine, as well as			
products and f	oods, not to mention the safety and efficacy			
of medical devices and <b>radiation-emitting devices</b> . Furthermore, imported abovementioned goods to USA from abroad is subject to the FDA's supervision.				
The scope of the FDA's regulatory	is very broad. The FDA's			
responsibilities are in close ties with several Department of Agriculture (USDA), the Drug and Border Protection (CBP), and the Consume	Enforcement Administration (DEA), Customs			
The following is a list, albeit not an exhaustive I	ist, of traditionally-recognized product			
that fall under t	the FDA's regulatory jurisdiction.			
Foods include:				
• dietary				

food additives

bottled

- infant formulas
- (other food products)\*

Drug	gs include:	
•	(brand-name + generic)	
•	non-prescription,	
Biolo	logics include: vaccines for humans	
•	blood and blood	products
•		gene therapy products
•	tissue and tissue component produ allergens	cts
Medi	dical devices include:	
•	simple items like tongue depressor	s and bedpans
•	complex	such as heart pacemakers
•	dental	
•	surgical	and prosthetics
Radia •	liation-emitting electronic products: microwave ovens	
•	x-ray	
•	laser products ultrasonic therapy equipment mercury vapor lamps sunlamps	

etc.

Cosme	etics include:			
•	colour additives found	d in makeup and othe	er	care
produ	cts			
•	skin	and cl	eansers	
•	nail	and pe	erfume	
Veteri •	inary products, such as livestock feeds pet foods	:		
•	veterinary		and devices	
• • •	co-based products, inco cigarettes cigarette tobacco roll-your-own tobacco e from the FDA, the US food products, such as	tobacco  Department of Agric		e in the regulating of
The F	DA also inspects		manufacturing	facilities worldwide,
includ	ing facilities that man	ufacture active		along with the
	ed product. Inspection d FDA staff. The FD		• •	
		from both the pu	ıblic and from the ind	ustry itself. The FDA
will d	often use such rep	orts to identify s	sites for which fur	ther inspection or
		is needed. The m	najority of companies	under inspection are
found	to be adequately com	npliant with the Cur	rent Good Manufactu	ring Practice (CGMP)

regulations.

The FDA is responsible for the approval and market surveillance of food, medicines and medical devices. As such, the authority is endowed with a police force and may pass laws within the limits specified in the Code of Federal Regulations (21 CFR)

Although the FDA has advantages of having centralization and common rules, there are several procedures stipulated under a 510(k) for Class II, or a PMA, or premarket approval, for Class III medical device that the EU regulates through a network of centralized and decentralized agencies throughout its member states.

The FDA is known and feared for its on-site	through the				
resulting deviation reports (i.e. Form 483) and warning letters, as it	monitors the market				
more actively than European authorities, searching for	*				
products and medical devices.					
The consequences of are more serio	ous. They range from				
publishing all the crimes to violent	closures, from				
bans to the withdrawal of all products	on the market by US				
marshalls—needless to say, at the company's expense.					

## Aufgabe Lückentext:

Folgende Wörter bitte in den Lückentext einfüllen. Jedes Wort kommt einmal vor. Bitte Gross- und Kleinbuchstaben beachten.

authority, biological, categories, cellular, component, devices, drugs, deviations, equipment, health, implants, ingredients, investigation, inspections, import, moisturizers, over-the-counter, polish, prescription, pharmaceutical, plant, personal, products, safety, supplements, smokeless, technologies, unwarranted, veterinary, water

<sup>\*</sup> They are those which may not be developed, manufactured and sold in accordance with the regulations.