Serotonin and norepinephrine reuptake inhibitors, selective serotonin reuptake inhibitors and other antidepressants and persistent sexual dysfunction - An EudraVigilance analysis

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Administrative details

EU PAS number	r
EUPAS39112	
Study ID	
39113	
DARWIN EU® study	
No	
Study countries	
United Kingdom	

Study description

This study was conducted to describe the case reports of persistent sexual dysfunctions to Serotonin and norepinephrine reuptake inhibitors, selective serotonin reuptake inhibitors and other antidepressants.

Study status

Finalised

Research institutions and networks

Institutions

European Medicines Agency (EMA)

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Institution

Contact details

Study institution contact

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Study contact

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Primary lead investigator

Pinheiro Luis

Primary lead investigator

Study timelines

Date when funding contract was signed

Planned: 04/12/2018 Actual: 07/12/2018

Study start date

Planned: 07/12/2018 Actual: 07/12/2018

Date of final study report

Planned: 13/02/2019 Actual: 26/02/2019

Sources of funding

EMA

Regulatory

Was the study required by a regulatory body?

Yes

Is the study required by a Risk Management Plan (RMP)?

Not applicable

Methodological aspects

Study type

Study topic: Human medicinal product Disease /health condition Study type: Non-interventional study Scope of the study: Other If 'other', further details on the scope of the study Case series review of Pharmacovigilance data **Data collection methods:** Secondary use of data Main study objective: Describe cases of Post-SSRI sexual dysfunction Study Design

Non-interventional study design

Other

Non-interventional study design, other

Case-series

Study drug and medical condition

Study drug International non-proprietary name (INN) or common name

CITALOPRAM

FLUOXETINE

FLUVOXAMINE

PAROXETINE

PAROXETINE ACETATE

SERTRALINE

DESVENLAFAXINE

FLUOXETINE HYDROCHLORIDE

PHENTERMINE HYDROCHLORIDE

OLANZAPINE

Medical condition to be studied

Loss of libido

Genital hypoaesthesia

Libido decreased

Female sexual arousal disorder

Anorgasmia

Female orgasmic disorder

Male orgasmic disorder

Additional medical condition(s)

Post-SSRI Sexual Dysfunctions

Population studied

Short description of the study population

Case reports of persistent sexual dysfunctions to Serotonin and norepinephrine reuptake inhibitors, selective serotonin reuptake inhibitors and other

Age groups

Children (2 to < 12 years)

Adolescents (12 to < 18 years)

Adults (18 to < 46 years)

Adults (46 to < 65 years)

Adults (65 to < 75 years)

Adults (75 to < 85 years)

Adults (85 years and over)

Estimated number of subjects

3210

Study design details

Data analysis plan

Descriptive statistics were run on the totality of the cases with sexual dysfunctions, by reporter type, by substance, by class, by indication, by country, by age, gender and by previous medical history of sexual dysfunction. Results by reaction were stratified by core PSSD case definition (see 4.3.2.), extended terms definition, which are the remaining terms from the sexual dysfunction case definition (see 4.3.1.) and secondary terms as per annex II.

Documents

Study results

Antidepressants and PSSD - EV analysis - report - 20190220.pdf(899.54 KB)

Data management

Data sources

Data source(s), other EudraVigilance	
Data sources (types)	
Spontaneous reports of suspected a	adverse drug reactions
Use of a Common Dat	a Model (CDM)
CDM mapping	
No	
Data quality specificat	tions
Check conformance	
Unknown	
Check completeness	
Unknown	
Check stability	
Unknown	
Check logical consistency	
Unknown	

Data characterisation

Data characterisation conducted

No