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**CONTENTS**

ETYOLOGY OF CIRCULAR CARIES .....	829
Olga Kokoceva-Ivanovska .....	829
Efka Zabokova-Bilbilovska .....	829
Mijoska Aneta.....	829
BIOCOMPATIBILITY AND REACTION OF DENTAL POLYMERS IN ORAL ENVIRONMENT .	835
Natasha Stavreva .....	835
Biljana Kapusevska .....	835
STUDY OF THE SIZE OF THE ANGLE OF THE MANDIBLE.....	841
Svetlana Jovevska.....	841
Mihajlo Petrovski .....	841
DENTAL SCANNERS IN PROSTHODONTICS .....	845
Blagoja Dashtevski.....	845
Aneta Mijoska.....	845
Marjan Petkov.....	845
Vanco Spirov .....	845
Oliver Dimitrovski.....	845
TEETH EXTRACTION WITH ROOT DELACERATION ON THE ORTHODONTIC INDICATIONS .....	853
Pavel Stanimirov.....	853
Greta Yordanova.....	853
DIFFERENCE IN CRANIOFACIAL CHARACTERISTICS IN ATTRACTIVE MALE AND FEMALE FACES .....	859
Sofija Carceva Shalja.....	859
Sandra Atanasova .....	859
Mihajlo Petrovski .....	859
ORAL TISSUE CONDITION IN PROSTHODONTIC GERIATRIC PATIENTS .....	865
Aneta Mijoska.....	865
Gordana Kovacevska .....	865
Georgi Tomov.....	865
INTRODUCTION TO ORTHODONTIC EDUCATION IN BULGARIA OF DIGITAL TECHNOLOGIES AND 3D ARCHIVING .....	869
Greta Yordanova.....	869
Martin Mladenov .....	869
INDICATIONS FOR MOLAR EXTRACTION DURING ORTHODONTIC TREATMENT .....	873
Sandra Atanasova .....	873
Sofija Carceva Salja.....	873
Ljubica Prosheva .....	873
COMPOSITE RESTORATIONS IN DENTISTRY .....	879
Aleksandar Andreevski.....	879
INFLUENCE OF GROWTH IN TREATED CLASS III GROWING PATIENTS WITH FACE MASK THERAPY AND UNTREATED PATIENTS.....	883
Sofija Carceva Shalja.....	883
Sandra Atanasova .....	883
POSITION OF THE MENTAL FORAMEN IN CORELATION WITH THE LOWER PREMOLARS: A PANORAMIC RADIOGRAPHIC STUDY .....	889
Mihajlo Petrovski .....	889
Svetlana Jovevska.....	889
Olivera Terzieva-Petrovska .....	889

PREVALENCE OF DENTAL CARIES IN CORRELATION WITH ORAL HYGIENE .....	895
Ivona Kovacevska.....	895
Natasha Longurova.....	895
Katerina Zlatanovska.....	895
THE SIGNIFICANCE OF TRAINING AND EDUCATION IN ACCURATE EVALUATION OF NATURAL TOOTH COLOR .....	901
Julija Zarkova .....	901
Ivan Nacevski .....	901
Vesna Korunovska Stefkovska .....	901
FUNCTIONAL RECOVERY OF VOICE FUNCTION IN PATIENTS WITH LARYNGEAL DYSFUNCTION: OBSERVATIONAL STUDY .....	905
Galina Mratskova .....	905
Damyan Petrov .....	905
THE INFLUENCE OF HYPERPROLACTINEMIA ON THE LEVEL OF FSH AND LH IN WOMEN .....	911
Mire Spasov.....	911
Verica Spasova .....	911
ACREDITED MOLECULAR METHODS FOR DETECTION OF INFECTIONS CAUSED BY HIV, HEPATITIS B VIRUS AND HEPATITIS C VIRUS .....	917
Belinda Gelmanovska.....	917
Vaso Taleski .....	917
FLUORESCENT IN SITU HYBRIDIZATION AND IMMUNOHISTOCHEMISTRY FOR SUBTYPING “NON-CLASSIFIABLE” RENAL CELL CARCINOMAS .....	921
Atanas Ivanov .....	921
Vili Stoyanova .....	921
JEJUNAL ADENOCARCINOMA: A CASE REPORT .....	925
Gordana Bozhinovska Beaka.....	925
Biljana Prgova Veljanovska .....	925
Milka Zdravkovska.....	925
Irena Eftimovska Rogac .....	925
Nadica Bozhinovska .....	925
THE ROLE OF TUMOR MARKERS CA 125, CA 72-4 AND CA 19-9 IN DETECTION AND MONITORING OF THE COURSE OF DISEASE OF OVARIAN CARCINOMA AND CARCINOMA OF THE UTERINE BODY .....	931
Kristina Petkova .....	931
Venci Chalkov .....	931
SPLENIC ABSCESS IN PATIENT WITH CHRONIC PANCREATITIS: A CASE REPORT .....	937
Gordana Bozhinovska Beaka.....	937
Biljana Noveska-Petrovska.....	937
Biljana Prgova Veljanovska .....	937
Nadica Bozhinovska .....	937
ROLE OF GOUT IN ATRIAL FIBRILLATION.....	943
Antoniya Kisheva .....	943
DEVELOPMENT ROLE AND IMPORTANCE OF FETAL AUTOPSY SCIENTIFIC PURPOSES - REVIEW OF THE LITERATURE.....	947
Tanya Kitova .....	947
BARRIERS TO THE EFFECTIVENESS OF DISEASE MANAGEMENT IN PEOPLE WITH DIABETES MELLITUS .....	953
Boryana Levterova .....	953

---

DYABETTE CONTROL PROGRAM AND PREVENTION OF THEIR CONDITIONS - PROPOSALS AND GUIDELINES .....	959
Varvara Pancheva .....	959
Valentin Vassilev .....	959
ALGORITHM OF COMPLEX PROGRAM FOR TREATMENT AND REHABILITATION IN DEGENERATIVE DISEASES OF THE VERTEBRAL COLUMN IN THE CERVICAL AREA. PREVENTION OF CERVICAL ARTHROPATHY .....	963
Petya Kasnakova .....	963
THE DIAGNOSIS OF DAUCUS CAROTA AS ALERGOGEN ON THE IMMUNE SYSTEM IN WHITE LABORATORY RAT .....	969
Mire Spasov .....	969
Icko Gjorgoski .....	969
THE DIAGNOSIS OF PHLEUM PRATENSE AS ALERGOGEN ON THE IMMUNE SYSTEM IN WHITE LABORATORY RAT .....	975
Hristijan Spasov .....	975
Mire Spasov .....	975
Icko Gjorgoski .....	975
Majlinda Ademi .....	975
INCREASED MEASLES MORBIDITY AND PUBLIC AWARENESS OF THE RISKS OF REFUSING IMMUNIZATION FOR THIS DISEASE .....	981
Ivelina Dobreva .....	981
Pavlina Teneva .....	981
Tsvetana Bojkova .....	981
PERSPECTIVES IN HEALTH CARE FOR CHILDREN WITH SPECIAL NEEDS IN CHILDREN'S AND SCHOOL HEALTH .....	985
Monika Obreykova .....	985
Galina Terzieva .....	985
MYOPIA IN PRE-SCHOOL CHILDREN AND MOBILE PHONES .....	991
Aleksandar Dodevski .....	991
Strahil Gazepov .....	991
Biljana Dodevska .....	991
Alen Georgijev .....	991
VISUAL SCREENING OF SCHOOL CHILDREN IN THE MUNICIPALITY OF SAPAREVA BANYA IN SOUTHWESTERN BULGARIA .....	995
Slavena Stoykova .....	995
Ekaterina Petrova .....	995
WETTABILITY – ANOTHER PARAMETER FOR GOOD FITTING OF A SILICONE-HYDROGEL DAILY DISPOSABLE CONTACT LENS .....	1001
Nikola Peev .....	1001
SUCCESSFUL COGNITIVE AGING - COGNITIVE RESERVE AND NEUROPLASTICITY .....	1005
Antonia Yaneva .....	1005
Kristina Kilova .....	1005
Teodora Dimcheva .....	1005
THE ROLE OF ACUPUNCTURE IN THE REHABILITATION OF PARALYSIS OF N.FACIALIS .....	1009
Lence Nikolovska .....	1009
Stefanija Gjorgieva .....	1009
Mire Spasov .....	1009

---

NEED FOR TRAINING OF HEALTHCARE PROFESSIONALS FROM OUTPATIENT CARE FOR BURNS INJURIES .....	1013
Anushka Dimitrova.....	1013
CARE AND TREATMENT IN PATIENTS WITH HEPATITIS B IN THE MUNICIPALITY OF PRILEP .....	1019
Panova Gordana.....	1019
Boshevska Viktorija .....	1019
Liljana Simonovska .....	1019
Iva Paneva .....	1019
Lenche Nikolovska .....	1019
Mire Spasov .....	1019
Gjorgji Shumanov.....	1019
ACUTE VENLAFAKINE OVERDOSE WITH POSITIVE URINE IMMUNOASSAY FOR TRAMADOL – CLINICAL AND DIAGNOSTIC OVERLAP - CASE REPORT AND LITERATURE OVERVIEW .....	1027
Pereska Zanina.....	1027
Janicevic-Ivanovska Danijela .....	1027
Bekjarovski Niko.....	1027
Simonovska Natasha.....	1027
Babulovska Aleksandra .....	1027
DRUG-INDUCED NEUTRALIZING ANTIBODIES TO TNF- $\alpha$ BLOCKES IN PATIENTS WITH INFLAMMATORY JOINT DISEASES FOLLOWED BY 24-MONTHS.....	1033
Krassimir Kraev.....	1033
Mariela Geneva-Popova .....	1033
Velichka Popova.....	1033
Stanislava Popova.....	1033
INFORMATION AND KNOWLEDGE OF THE ANTIBIOTICS AND THE ANTIBIOTIC RESIDENCE OF THE POPULATION FROM THE SOUTHEAST REGION OF REPUBLIC OF NORTH MACEDONIA .....	1039
Milka Zdravkovska.....	1039
Marija Darkovska-Serafimovska .....	1039
Gordana Bozinovska-Beaka .....	1039
Svetlana Zivkova .....	1039
SIMPLE LIQUID CHROMATOGRAPHY METHOD WITH UV DETECTION FOR DETERMINATION OF BROMAZEPAM IN SOLID PHARMACEUTICAL DOSAGE FORMS ....	1045
Irena Brčina .....	1045
Marija Darkovska Serafimovska .....	1045
Tijana Serafimovska .....	1045
Trajan Balkanov .....	1045
Biljana Gjorgeska .....	1045
CONSUMPTION ANALYSIS OF MOST PRESCRIBED ANTIBIOTICS FINANCED BY THE HEALTH INSURANCE FUND IN REPUBLIC OF NORTH MACEDONIA .....	1051
Elena Drakalska.....	1051
Bistra Angelovska.....	1051
Veneta Zdravkova.....	1051
MICROBIOLOGICAL DIAGNOSIS AND IMPORTANCE OF INFECTIONS CAUSED BY INFLENZAE – A VIRUS .....	1057
Zivadinka Cvetanovska .....	1057
Vaso Taleski .....	1057

---

POISONOUS SNAKES OF BULGARIA: CLINICAL AND EPIDEMIOLOGICAL ANALYSIS .....	1063
Olimpiada Atmazhova .....	1063
Evgenia Barzashka .....	1063
Iskra Petkova .....	1063
BODY FAT DISTRIBUTION AND LIPID PROFILE CHANGES AFTER WEIGHT LOSS – A CASE REPORT .....	1071
Slavica Shubeska Stratrova .....	1071
Danijela Janicevic Ivanovska .....	1071
FACTORS FOR OBESITY IN STUDENTS FROM THE SCHOOLS .....	1077
Antoaneta Grozeva .....	1077
INFLUENCE OF SUNSHINE ON THE SERUM LEVELS OF VITAMIN D .....	1081
Ivelina Dobreva .....	1081
Pavlina Teneva .....	1081
Valya Trencheva..	1081
GEOMEDICAL TOURISM – OUTDOOR HEALTH CARE FACILITIES ANTI-GEOPATHIC STRESS TREATMENT .....	1087
Krasimira Staneva.....	1087
APPLICATION OF BLASTWARE SOFTWARE FOR MEASURING MICROCOLIMIC CONDITIONS .....	1093
Naim Baftiu .....	1093
Raif Bytyqi .....	1093
Overall, the number of measurements .....	1096
Percentage .....	1096
APPLICATION OF BIOPRINTING IN CONTEMPORARY MEDICINE (REVIEW).....	1099
Desislava Bakova .....	1099
Kristina Kilova .....	1099
Maria Semerdjieva.....	1099
CBRN EVENT - DO WE NEED MEDICAL INTELLIGENCE .....	1103
Elena Valkanova.....	1103
Rostislav Kostadinov .....	1103
ACUTE PANCREATITIS .....	1107
Redzep Emurlai .....	1107
PALLIATIVE CARE AND EUTHANASIA – CONVERGENCE POINTS AND PUBLIC ATTITUDES .....	1113
Nikola Sabev.....	1113
SOCIOECONOMIC CHARACTERISTICS OF PATIENTS WITH TENSION HEADACHE IN MUNICIPALITY OF VELES .....	1119
Elena Manchevska.....	1119
Toshe Krstev.....	1119
Gordana Panova.....	1119
SEASONALITY IN THE APPEARANCE OF INTRA HOSPITAL INFECTIONS .....	1127
Emilija Chapevska .....	1127
PEDAGOGICAL COMPETENCE OF MENTORS DURING THE PRE-GRADUATION TRAINEESHIP OF FUTURE NURSES .....	1131
Snezhana Dragusheva.....	1131
Penka Petleshkova .....	1131
Tanya Paskaleva .....	1131

**ACUTE VENLAFAXINE OVERDOSE WITH POSITIVE URINE IMMUNOASSAY FOR TRAMADOL – CLINICAL AND DIAGNOSTIC OVERLAP - CASE REPORT AND LITERATURE OVERVIEW**

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**Abstract:** Objective. The overlapping of pharmacokinetics and/or the pharmacodynamics of medicines causes the occurrence of overlapping clinical syndromes and diagnostic issues, potentiated in overdoses. We report a case of severe venlafaxine poisoning where the clinical presentation and the results of rapid immunoassay test overlapped with tramadol intoxication.

Case presentation. An unconscious women with recurrent seizers, hypertension and supposed acute medication poisoning in suicidal attempt was transported to our clinic. Previously, she had been lavaged, rehydrated and treated with 20 mg diazepam iv, 40 mg furosemide at the local general hospital. Her regular tablet therapy consisted of losartan, levothyroxine, venlafaxine, occasionally tramadol.

At admission she was comatose, with isochoric normal pupils, BP 130/80 mm Hg, SaO<sub>2</sub> 86%, and recurrent episodes of seizures treated with 10mg diazepam iv, ocular clonus, hypertonus, temperature 38.9C, diaphoresis, facial hyperaemia, dark coloured urine, hyponatremia and rhabdomyolisis. The lateral flow immunoassay (AbuGnostR) was positive for tramadol, but the homogeneous enzyme immunoassay did not confirm it. After 36 hours of intensive treatment she became somnolent and reported ingestion of 2250 mg tbl Venlafaxine. The AbuGnost R test detects tramadol at cut off urine values 200ng/ml, but present cross reactivity with O-desmethyl-venlafaxine at cut off values up to 25000ng/ml. The following days she complained of muscular weakness, headaches and cognitive impairment, which lasted for more then one month after release from hospital.

Conclusion. High concentrations of venlafaxine metabolites induce false positive tramadol immunoassay (AbuGnostR) test. Overlapping clinical presentations and metabolic pathways of venlafaxine and tramadol should alert physicians when interpret rapid immunoassay test. The mandatory principle when making medical decisions should cover synthesis of critically interpreted toxicology analysis, interview data and clinical features of the poisoning, which may help to avoid misleading conclusions and improve the diagnostic and therapy decisions.

**Keywords:** venlafaxine, tramadol, poisoning, lateral flow immunoassay,

## 1. INTRODUCTION

Venlafaxine is an antidepressant acting as a serotonin and norepinephrine reuptake inhibitor and weak inhibitor of dopamine reuptake. The reported poisonings with venlafaxine showed lower toxicity compared to TCA, but higher than SSRIs. The complexity of treatment decisions-making for these poisonings is associated not only with the disproportionate poisoning severity to the ingestion dose (Fischer, Unterecker, & Pfuhlmann, 2012) and overlapping clinical syndromes with other serotonin and/or norepinephrine reuptake inhibitors, but also with the necessity for rapid diagnosis in clinical practice and a high likelihood of false positives of rapid immunoassay tests (Saitman, Park, & Fitzgerald, 2014).

We report a case of severe venlafaxine poisoning which clinical presentation and rapid immunoassay test (LFIA-lateral flow immunoassay) overlapped with tramadol intoxication.

## 2. CASE REPORT

A middle aged women was admitted at the clinic after two prolonged episodes of seizures and supposed acute poisoning with medicines in suicidal attempt. She had a history of hypothyreosis, regularly treated with tablet (tbl) euthyrox (75 mg/day), hypertension treated with tbl losartan (50mg/day), depression treated with tbl. venlafaxine 75 mg/day and traumatic injury of the right ankle and foot treated with capsule tramadol a 50 mg occasionally. After her family found her unconscious, she was taken to the local general hospital where gastric lavage was performed and crystalloids were administered. After half an hour she had seizures lasting a couple of minutes. The second episode of seizures was associated with dramatic increase of blood pressure so 10 mg diazepam intravenously (iv) with 40 mg furosemide intramuscularly (im) were administered and she was transferred to our Clinic. During the transportation she received additional 10 mg diazepam im. because of new episode of seizures.

At admission she was comatose, with isochoric pupils, BP 130/80 mm Hg, SaO<sub>2</sub> 86% with clear lungs, soft abdomen and electrocardiogram (ECG) with sinus rhythm HR 95/min, QRS 105msec, QTc 437msec, non ischemic (ascendant) ST elevation.

Soon, she had another episode of seizures associated with more severe hypoxemia (75%) treated with diazepam 10 mg iv and gradually she developed ocular clonus, with hypertonus, increased temperature 38.9C and diaphoresis with facial hyperaemia. Urinary catheter insertion revealed dark urine.

Laboratory findings showed increased white blood counts, creatine kinase (CK), myoglobin and severe hyponatremia 111mmol/l (137-145 mmol/l referent values). Serum analysis: BUN, creatinine, glycemia, bilirubinemia, alkaline phosphatase, gamma-glutamyltransferase, calcium were in referent range during the whole hospital stay of the patient (Table 1).

**Table 1. Laboratory findings during hospital stay**

Parameter	Day 1	Day2	Day3	Day4	Day6	Day 8	Referent
Le x109/l	23.6	17,3	10,6	10,4	6,3	7,1	4-11
AST (U/l)	31	32	38		67	29	10-34
ALT (U/l)	20	19	22		67	46	10-45
LDH (U/l)	392	276	254		223	209	<248
Na (mmol/l)	109	120	135		137	136	137-145
K (mmol/l)	3,6	3,6	3,4		3,4	3,9	3,8-5,5
CRP (mg/l)	0,6	29,9	68,1		7,1	1,9	<6
CK (U/l)	452	525		4234	1409		24-173
CK-MB (U/l)	33			40			<25
Myoglobin (ng/ml)	351	/	277	601	227	57	10-46
Troponin (ng/l)	1,3	/	7,3	6,1	/	/	<15,6

AST-Aspartate aminotransferase, ALT- Alanine aminotransferase, LDH-Lactate dehydrogenas, CRP-C-reactive protein, CK-Creatine kinase, CK-MB- Cretaine kinase-MB

The urine screening test (LFIA-AbuGnostR-BioGnost) for tramadol was positive on initial presentation. The patient was treated with crystalloids (0,9% NaCl) with dose regime 2.5 mL/kg/h IV infusion, hypertonic sodium (according Joint European Guidelines), sodium hydrogen bicarbonate (1mmol/kg i.v ), diazepam 0.20mg/kg iv, ceftriaxone 2gr/day.

After 36 hours of intensive therapy and after serum concentrations of sodium were normalised, the seizures ceased. The patient was somnolent, complained of muscular weakness, difficulties in self-standing from bed and upright standing, spasm of jaw muscles, headaches, with cognitive difficulties and disorientation. She self-reported ingestion of 30 tbl venlafaxine 75mg. The re-analysis of the firstly collected urinary sample using homogenous enzyme immunoassay HEIA (Beckman coulter AU 480) excluded the presence of tramadol and its metabolites in the patient's urine. The following days gradual normalization of consciousness occurred but the headaches and cognitive deterioration persisted. Plasma levels of myoglobin reached their maximum on the fourth day of hospitalization with parallel existence of expressed muscle weakness in the same period. Treatment with intensive hydration, bicarbonates and diuretics enabled the preservation of renal function.

Consultative examination by the neurologist on the 4th day confirmed that there were no neither focal neurologic deficit nor meningeal irritation, but also noted that the patient was confused, disoriented with difficulties in maintaining contact. During the hospital stay she presented slow regression of the parameters of rhabdomyolysis with preserved renal function, and persistent cognitive impairment.

ECG at discharge presented sinus rhythm, HR 95/min, normal P wave, QRS 85msec, QTc 407msec, isoelectric ST segment.

A month after release from hospital, due to persistent headaches, selective amnesia, confusion, anxiety and insomnia, the patient was re-examined by a neurologist who excluded organic neurological suffering and deficit. She continued the treatment at the psychiatric unit.

### **3. DISCUSSION**

There were several reports of cross-reactivity of venlafaxine with phencyclidine using rapid immunoassays(Bond, Steele, & Uges, 2003) but our presentation was first reporting of venlafaxine with tramadol cross reactivity using rapid LFIA test in acute venlafaxine overdose (VO). The goal of presenting this case is to provide a wider clinical context when interprets urine drug test results considering overlapping clinical presentation of venlafaxine poisoning with tramadol and reassessment of the inefficacy if antidote treatment (naloxone) was used in such poisonings.

Venlafaxine (antidepressant) and tramadol (analgesic) are prescribed for different medical indication but share many similarities in chemical structure, inhibition of serotonin and nor-adrenalin reuptake and metabolic pathway(Reeves & Cox, 2008). Even more, it was found that venlafaxine present analgesic effects(Jha, Mazumdar, & Bhatt, 2006) as well as tramadol have antidepressant activity(Kalra, Tayal, & Chawla, 2008). In our case, the patient was receiving concomitant treatment with tbl venlafaxine and occasionally cap. tramadol, according medical recommendation. Poisonings with venlafaxine as well as with tramadol induce overlapping clinical features, associated with their pharmacokinetic profile.

Rhabdomyolysis is a significant clinical feature of acute poisoning with antidepressants, especially with venlafaxine(Wilson, Howell, & Waring, 2007), but it is associated with acute tramadol poisoning too(Khan, Yousef, & Errayes, 2010). The laboratory findings confirmed the development of rhabdomyolysis with more than five time increase of CK (creatine phosphokinase) at the 4th day of hospitalisation and serum myoglobin as an early marker of myotoxicity (Melli, Chaudhry, & Cornblath, 2005). Myoglobinuria was not investigated in our patient due to the unavailable diagnostic tests, although some studies have found that the absence of urine myoglobin does not exclude rhabdomyolysis. Although, seizures are confounding factor for developing of rhabdomyolysis, the proposed underlying mechanism is direct myotoxic effects of venlafaxine6, and also of tramadol poisoning(Rahimi, Soltaninejad, & Shadnia, 2014); the cardiac source of increased CK and myoglobin was excluded with ECG and normal troponin serum concentrations in our patient.

Hyponatremia (serum Na <135 mmol/l) is a serious electrolyte disturbance associated with poisoning with new generation of antidepressants including venlafaxine and tramadol poisoning(De Picker, Van Den Eede, Dumont, Moorkens, & Sabbe, 2014) too. Patients under diuretic therapy and elderly ones are at higher risk for hyponatremia, even in regular pharmacology doses (De Picker et al., 2014). The proposed mechanism is associated with serotonin reuptake inhibition, inducing either increased release of antidiuretic hormone (ADH) or increased sensitivity to ADH implying on syndrome of inappropriate secretion of ADH(De Picker et al., 2014). VO is associated with prominent hyponatremia(De Picker et al., 2014), even more pronounced than the poisonings with other types of SSRI. Our patient presented severe hyponatremia (<115 mmol/l) according Joint European Guidelines with life - threatening neurologic features (seizures, coma). Seizures cessation after correction of hyponatremia in our case was a clinical confirmation of the electrolyte imbalance as an underlying mechanism.

Serotonin syndrome (SS) in venlafaxine overdose is well documented in literature(Abadie et al., 2015). Although there was a suspicion that a tramadol poisonings causes SS(Ryan & Isbister, 2015), more studies have shown that tramadol overdose also induces the onset of SS (Tashakori & Afshari, 2010). SS is potentially life -threatening condition, confirmed in our patient according Hunter diagnostic criteria(Dunkley, Isbister, Sibbritt, Dawson, & Whyte, 2003). After complete clinical and laboratory restitution, the patient suffered from prolonged neurologic impairment, already reported in the literature considering VO.

The overlapping pharmacokinetic profile of venlafaxine and tramadol generates false positive results in diagnostic toxicological tests, too. The false positive result for tramadol LFIA in our patient resulted from the venlafaxine metabolite, O-desmethyl-venlafaxine which has similar transition to tramadol (O-desmethyl-tramadol) inducing cross-reactivity (Allen, 2006). The HEIA test from the same urine sample was negative for tramadol because has no cross-reactivity with venlafaxine and its metabolite O- desmethyl-venlafaxine; together with self-reported venlafaxine overdose, leaded to conclusion that the positive rapid LFIA test in fact diagnosed the venlafaxine poisoning. The AbuGnost R test detects tramadol with urine cut off values 200ng/ml, but present cross reactivity with O-desmethyl-venlafaxine at cut off values up to 25000ng/ml.

We presented severe VO with self-reported ingested dose of 2250 mg. Most of the cases reported association of fatal outcome with ingested venlafaxine doses in tens of grams as a contrary to the case report of poisoning with 3 grams

venlafaxine with lethal outcome where venlafaxine plasma concentrations were verified by LC-MS/MS (Vignali, Morini, Chen, Stramesi, & Groppi, 2014). The differences in the venlafaxine metabolism is associated with CYP2D6 polymorphism which induce poor, intermediate, extensive and ultrarapid venlafaxine metabolizing phenotypes(Rolla et al., 2014). These facts should be considered when observing poisonings with ingested lower venlafaxine quantities and pay attention to the clinical signs of SS and laboratory findings.

The ingested venlafaxine dose was presented based only on patient's self reporting due to the unavailability of serum venlafaxine quantification, which is a limitations of our analysis. However, the positive data from the interview, the course of clinical presentation, the results from laboratory analysis (hyponatremia), the exclusion of tramadol as the causative agent of poisoning with toxicological analyzes and prolonged patient's cognitive deterioration, were sufficient to confirm the venlafaxine poisoning. Although the clinical presentation of venlafaxine and tramadol as well as their symptomatic treatment overlap to a large extent, the dilemma about absence of clinical improvement when use the antidotal therapy (such as naloxone) in the event of respiratory failure is a key point when treating these two types of poisonings only according to the results of the rapid tests.

#### **4. CONCLUSION**

The role of rapid immunoassay test is significant and necessary for the work of the emergency departments, taking in consideration that high concentrations of venlafaxine metabolites induce false positive tramadol immunoassay (AbuGnostRBognost) test and overlapping clinical presentations. The mandatory principle when making medical decisions should cover synthesis of critically interpreted toxicology analysis, interview data and clinical features of the poisoning, which may help to avoid misleading conclusions and improve the diagnostic and therapy decisions.

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