Life Threatening Acute Heart Failure in Two Young Adults Treated with Antidepressant Medication

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Abstract

Duloxetine is a selective serotonin (5-HT) and norepinephrine reuptake inhibitor (SNRI) that is mainly prescribed for major depressive disorder and generalized anxiety disorder. It is our clinical experience that SNRI medication has caused severe heart failure in several patients, including one treated with Impella (LVAD).

Here we present two cases of young adults treated with Duloxetine who developed acute heart failure but had different outcomes.

Acute heart failure following treatment is not a registered side effect of SNRI medication. Our cases and other published case reports suggest that noradrenergic reuptake inhibitors like Duloxetine can cause catecholamine surges and result in reversible cardiomyopathies if no further complications occur.

Keywords: Acute heart failure, Antidepressants, Duloxetine, Left ventricular assist

Introduction

Duloxetine is a selective serotonin (5-HT) and norepinephrine reuptake inhibitor (SNRI). It is approved in the United States for major depressive disorder, generalized anxiety disorder, diabetic peripheral neuropathic pain, fibromyalgia, and chronic musculoskeletal pain. Its mechanism of action is by inhibiting the reuptake of serotonin and norepinephrine thereby increasing their concentrations in different locations in the central nervous system. It further causes a weak inhibition of the reuptake of dopamine with a negligible affinity for adrenergic, cholinergic, dopaminergic and histamine receptors. Stress cardiomyopathy has been reported after catecholamine administration [1], and has been associated to patients with pheochromocytoma [2,3]. One case report suggests that adrenergic medications like Duloxetine can cause catecholamine surges and result in reversible cardiomyopathies [4].

Acute left ventricular (LV) dysfunction, e.g. Takotsubo cardiomyopathy or stress cardiomyopathy, usually follows sudden emotional or physical stress [5]. The mechanism is believed to be mediated through epinephrine and causes stunning of the myocardium, with a predilection for the cardiac apex [6].

Current lists of side effects of Duloxetine do not enlist cardiomyopathy but do entail shortness of breath [7]. Patients treated with Duloxetine are not routinely followed for signs of heart failure.

Methods

Two individuals were admitted to our cardiology unit at Uppsala University Hospital with acute heart failure. The authors were clinically involved and responsible for the initial care and follow-up for both patients. Reports were reviewed and approved by the local Ethics Committee and both patients provided written consent forms for the publication.

Case 1

A 36-year-old female with a history of depression and anxiety disorder had been medicated with Duloxetine 60 mg daily for 18 months and Mirtazapin for 6 months. For a period of six months she experienced increased lethargy and a 20 kg weight gain. Her symptoms were initially perceived as aggravated depressive symptoms.

At her local hospital she was diagnosed with dilated cardiomyopathy with a reduced left ventricular ejection fraction (LV-EF) of 20-25%. Her coronary angiography was normal. A cardiac MRI was inconclusive, i.e. had no definite sign of an earlier myocarditis. Her liver function tests (LFTs) were elevated. Heart failure management was initialized and she was planned for regular follow-up.

Two months later her heart failure symptoms progressed to NYHA class IV. She was dehydrated with increased creatinine 110 μmol/L (baseline 80 μmol/L), hemoglobin was 140 g/L (baseline 120 g/L), CRP 67 mg/L and NT-proBNP levels of 5800 ng/L. Her LFTs were further elevated. Transthoracic echocardiography (TTE) showed a dilated left ventricle (LV) of 7 cm, moderately dilated right ventricle (RV), a moderate to severe mitral insufficiency and severely reduced LV-EF of 10-15%. Further, she had a thrombus located in the LV and an akinetic apex. Her systolic blood pressure was 80 mmHg but was stabilized with inotropes and she also received Levosimendan. Duloxetine and Mirtazapine were terminated and she received Warfarin. She was referred to the cardiology unit at Uppsala University Hospital.
Right heart catheterization (RHC) showed low pressures and low saturation in the pulmonary artery of approximately 40%, a cardiac output of 2.4 L/min, a cardiac index of 1.1 and an AV02-difference of 89%. She received Levosimendan again and received a Heart-Mate as a bridge to transplantation. Two months after the Heart-Mate operation her TTE showed LVEDD of 6.1 cm, normal sized RV but she continued to have severely reduced LV-EF of 15% and wide-spread hypokinesia. Postoperatively PA-pressures increased and she was treated with Revatio. She is currently working half-time and is waiting for a heart transplant.

Case 2

A 40-year-old obese female with a history of hypertension, migraine, depression and chronic back pain had been taking Duloxetine 90 mg daily for three years. For several months she had been under severe psychological stress because of a difficult domestic situation. Having been on benefits due to chronic back pain she was shortly before admission ordered to start a rehabilitation program, causing more stress. Further, she had been suffering from an upper respiratory tract infection for 3 weeks. Her other medications were Acetaminophen, Tramadol, Sumatriptan and Lactulose as needed.

She presented at the emergency room with sudden-onset heart failure, blood pressure 213/108 mmHg, tachycardia and loss of consciousness. Electrocardiogram (EKG) showed a regular sinus rhythm of 145 beats per minute and a left bundle branch block (LBBB). Hearth sounds were normal but she had bilateral crackles over both lungs. A TTE showed a dilated right and left ventricle and universal dyskinesia with a LV-EF of 35%. Laboratory tests showed BNP of approximately 1200 ng/L, CRP 120 mg/L, D-dimer of 12 and Troponin 764 ng/L. Coronary angiography was normal. CT thorax showed bilateral pleural effusion, no pulmonary embolus but pneumonia could not be excluded.

She was connected to a ventilator. Her systolic blood pressure dropped to 60 mmHg and she received low doses of Norpinephrine and Milrinone. She had intermittent VT attacks. Cardiac output measured by thermodilution was approximately 8 L/min and Sv02 80% and diuresis was normal.

Duloxetine was terminated on admission. She received a betablocker, an angiotensin receptor blocker and antibiotics. Five days later she was extubated and was fully alert. A new TTE revealed a normal LV-EF but she had a slight diastolic dysfunction with E/A of 0.9 and E/e’ of 11. BNP returned to normal and she no longer had a LBBB. She is currently in NYHA class I and is working full-time.

Discussion

Duloxetine is believed to have the highest relative norepinephrine selectivity of the SNRI medications [8]. In recent years a handful of case reports have been published suggesting causation between SNRI medication and heart failure [9-11]. Fangio et al. reported a case of venlafaxine (another norepinephrine reuptake inhibitor) poisoning complicated by acute heart failure with reduced systolic function, elevation of NT-proBNP and hemodynamic instability in a previously healthy young adult. The cardiac function was fully restored in ten days after termination of the SNRI medication [11]. The same authors state that the findings in their case report share certain features of Takotsubo cardiomyopathy. Christoph et al. described a case of Takotsubo cardiomyopathy in a patient after an accidental overdose of Venlafaxine [10,11]. According to our clinical experience Venlafaxine has caused severe heart failure in several patients, including one treated with Impella (LVAD) [12].

Natural disasters have been shown to increase the incidence of stress cardiomyopathy [13]. Other types of physical illness, i.e. intracranial bleeding, can cause surges of sympathetic response resulting in acute ECG changes in around 10% of cases, elevated cardiac enzymes and acute but reversible LV impairment in the presence of normal coronary arteries [14,15]. A profound increase in plasma catecholamine levels has been reported in many cases following stress cardiomyopathy and they have been shown to be up to 34 times higher than normal resting values and significantly higher than with than among patients with ACS or heart failure [16,17].

When epinephrine circulates at physiological and elevated concentrations it triggers an intracellular signaling switch through β2-adrenoceptors on cardiomyocytes of the ventricles causing a positive inotropic response. With high and long-standing (supraphysiological) epinephrine concentrations, however, they cause a negative inotropic effect on cardiomyocyte contraction [6]. This may cause myocardial apoptosis and heart failure because of LV dilatation [6]. One study has demonstrated that there is a higher concentration of β2-adrenoceptors in the apical myocardium in canine hearts [18]. If it is similar in humans, it may account for the apical predilection observed in most cases. The human ventricular myocardium has a higher concentration of β2-adrenoceptors than other mammals [19].

The use of norepinephrine reuptake inhibitors can increase the risk for heart failure with BNP elevation [16]. The apical wall motion abnormalities in stress cardiomyopathy are usually reversible within days or weeks of the acute insult (as is the case for traditional myocardial stunning) if no further complications occur [20]. A meta-analysis of forty-two placebo-controlled trials on individuals treated with Duloxetine showed a statistically significant increase in palpitations, tachycardia and hypertension [21].

Venlafaxine is mostly metabolized by the isoenzyme CYP2D6 and to a lesser extent by CYP3A4 [22]. Duloxetine is metabolized by CYP1A2 as well as CYP2D6. The pharmacokinetics of Duloxetine demonstrates great interindividual variability (usually around 50-60%) due to factors, e.g. sex, age, smoking status and ability to metabolize by means of the enzyme CYP2D6 [23]. For several psychotropics drugs, the CYP2D6 isoenzyme is a high-affinity/low-capacity enzyme whose polymorphisms can phenotypically determine slow, extended, or rapid metabolism [24].

We have presented two case reports of young adults treated with Duloxetine that resulted in life threatening acute dilated cardiomyopathy with different outcomes. Case 1 had been treated for 18 months with no apparent increase in the amount of stressors. Case 2 had taken Duloxetine for three years. The question arises why the acute decompensated heart failure occurred after such prolonged treatment.

The level of stress in Case 1 was relatively stable. Ferreira et al. reported a similar case of a 35-year old female treated with Venlafaxine for 3 months that presented with reversible synchronous pneumonitis and acute cardiomyopathy attributed to Venlafaxine [25]. We do not know whether Case 1 had a CYP2D6 slow metabolizer genotype and can only hypothesize whether she had drug accumulation and subsequent cellular/organic insult, either due to drug accumulation or concomitant surges of catecholamines. It is, however, an important fact that Duloxetine was not terminated until...
two months after Case 1 presented with heart failure symptoms. The delay may have caused the heart failure to be irreversible.

For Case 2 additional stressors were not present at the time she started the medication. Shortly before her decompensation she had found the additional stress to be extremely encumbering. We believe that the catecholamine surges caused by Duloxetine and increased stressors resulted in reversible cardiomyopathy. The cardiomyopathy, perhaps being a Takosubo type, is reversible if no further complications occur.

There are certain shortcomings to this paper. We did not perform a twenty-four hour urine collection of catecholamines. When we asked for titers of catecholamines in the blood test of our patients the blood tests had been disposed of. We did not have information on the intake of other supplements that may have inhibited the P450 enzyme complex leading to drug accumulation.

Conclusion

We believe that Duloxetine played a central role in potentiating severe stress cardiomyopathy in both patients. We suggest that certain precautions with regular ECG’s, BNP measurements and even annual TTE may be rationalized for patients treated with norepinephrine reuptake inhibitors. In case of heart failure symptoms we recommend that SNRI medication should be terminated.

Authors Agreement

No conflict to disclose.

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References

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