| 2 | | Coronary spasm as a cause of recurrent chest pain: a case report |
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| 4 5 | | Abstract |
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| 7 | | Background: |
| 8 | | Coronary artery spasm plays an important role in the pathogenesis of a wide variety of |
| 9 | Ì | ischemic heart diseases, including myocardial infarction and sudden cardiac death |
| 10 | Į | However, the diagnosis of vasospastic angina is not always easy on the basis of symptoms |
| 11 | | alone |
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| 12 | | Case presentation: |
| 13 | | A 36 year- old- man was evaluated due to recurrent left sided chest pain of 8 months |
| 14 | | duration. He is a smoker-, and otherwise physical examination was unremarkable. |
| 15 | | Electrocardiogram and blood tests were normal-, apart from elevated serum concentrations |
| 16 | | of triglycerides and low density lipoprotein cholesterol During the recovery phase of |
| 17 | | exercise treadmill test, the patient developed -progressive ST-segment elevation at inferior |
| 18 | I | leads II, III and aVF together with progressive ST depression at precordial leads V1 to V5. |
| 19 | | The ECG showed gradually decreased heart rate with the development of sinus bradycardia |
| 20 | | that progressed rapidly to complete heart block. The patient collapsed with chest pain and |
| 21 | | hypotension. It took about 10 minutes for the ECG retuning to the baseline after medical |
| 22 | | management. The patient was referred to the coronary care unit for further management |
| 23 | Ì | and underwent coronary angiogram. Coronary spasm -was observed in the proximal |
| 24 | ı | segment of the right coronary artery. The patient received medical treatment after coronary |
| 25 | | angiogram and stood well for more than one year follow up. |
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Conclusion

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- 27 Coronary spasm may lead to both recurrent chest pain and significant arrhythmia and the
- diagnosis often requires high index of suspicion and lab documentation.

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Background:

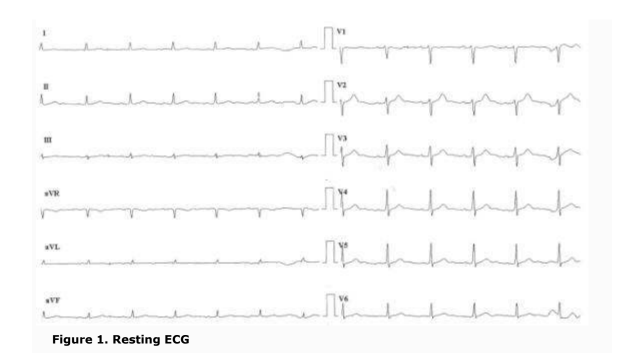
- 31 Coronary artery vasospasm, or smooth muscle constriction of the coronary
- artery, is an important cause of chest pain syndromes that can lead to
- myocardial infarction-, ventricular arrhythmias, and sudden death. Although it
- can occur in vessels distressed by atherosclerosis, traditionally it has been
- associated with variant or Prinzmetal's angina-, which was first described in
- 1959 (1). The diagnosis of vasospastic angina (VSA) is not always easy on
- 37 the basis of symptoms alone and often requires high index of suspicion and
- 38 lab documentation as well.

Case presentation:

- 40 A 36 year -old- man was evaluated at our out-patient clinic complaining of
- recurrent left sided chest pain of 8 months duration. The pain occurred at rest
- and was precipitated sometimes with effort-, which could be lasted for few
- 43 minutes-, diffuse-, vague in nature but not referred. The patient is a current
- 44 smoker for more than 10 years and did not have past history of diabetes-,
- 45 hypertension-, dyslipidemia or family history of coronary artery disease
- 46 (CAD).
- Physical examination was unremarkable with BMI at 24 kg/m²-, waist
- circumference at 95 cm-, and blood pressure was 140 /85 mmHg.

- Chest x-ray and resting electrocardiogram (ECG) were normal and blood tests
- showed low density lipoprotein cholesterol 3.99 mmol/L, high density
- 51 lipoprotein cholesterol 1.1 mmol/L,total cholesterol 5.72 mmol/L,triglycerides
- 3.13 mmol/L ,and HbA1c 5.0%. All other blood tests were normal.
- We decided to perform exercise stress test using CAEP protocol (The
- Chronotropic Assessment Exercise Protocol) (2). Blood pressure, heart rate
- 55 (HR) and 12-leads ECG were recorded at rest, at two-minute intervals during
- exercise, at peak exercise, and through the recovery phase. The ECG was
- continuously displayed and ST-segment was measured automatically by a
- computer-assisted system in all 12 leads. We decided to stop the test because
- the patient got fatigue with achievement of 89 % of age-predicted maximal
- 60 HR for age. No significant hemodynamic abnormalities or chest pain occurred
- with rapid upsloping ST-segment depression seen at maximal exercise.
- Achieved METs was 12.1 and RPP was 27 710 beats x mmHg. At minute 2
- in the recovery ,we noticed early ST-segment elevation in the inferior leads .
- HR was 122 beats/min, and then HR inappropriately decreased with
- progressive ST- segment elevation in the inferior leads together with
- progressive ST depression in the precordial leads V1 to V5. The patient
- started to get chest pain and feels dizzy. The ECG showed sinus bradycardia
- with 1st degree heart block followed by Mobiz-type II heart block and then
- complete heart block (CHB). It took about 4 min from cessation of exercise to
- develop CHB. HR was 30 beats/min, and BP was 65 /30 mmHg. At the start
- 71 ,the patient received oxygen and sublingual NTG . After the development of
- 72 CHB, he received 1 mg atropine iv push, and started fluid resuscitation. A
- random blood sugar was normal .After that by about 7 min, the ECG showed
- 74 junctional escape rhythm with HR 45 beats/min ,followed rapidly by

accelerated junctional rhythm with HR 101 beats/min then sinus tachycardia. Also, ST-segment changes in the inferior leads and precordial leads gradually improved till complete resolution after about 10 minutes from its start ,together with disappearance of chest pain and normalization of BP.



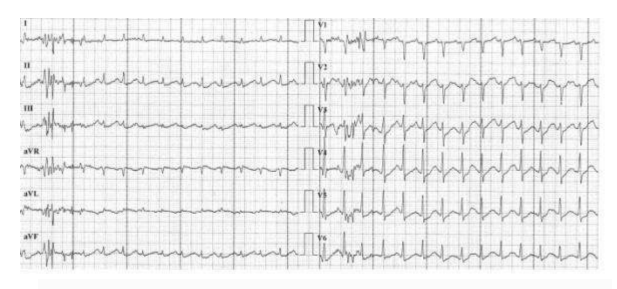


Figure 2. Maximal exercise treadmill test with normal response

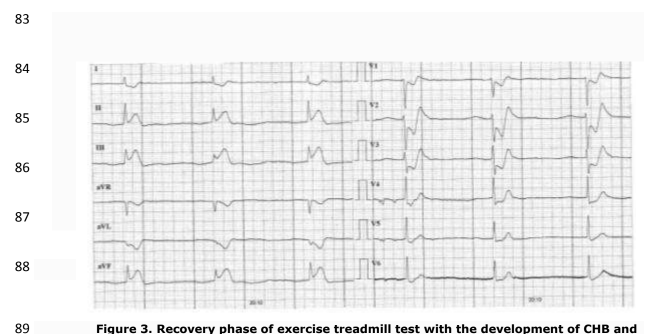
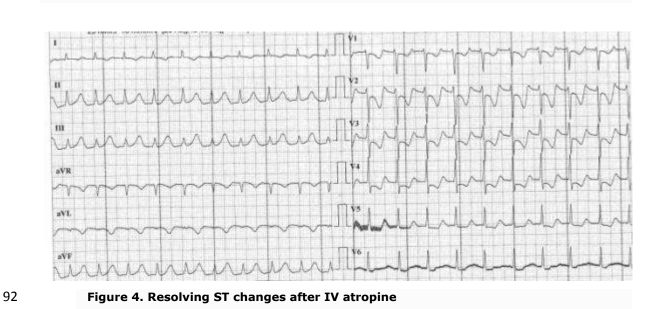


Figure 3. Recovery phase of exercise treadmill test with the development of CHB and ST elevation at inferior leads together with ST depression at precordial leads



The patient was transferred to the coronary care unit for observation and further evaluation. Cardiac enzymes were checked at baseline, 6, and 12 hours, and echo-doppler evaluation was done where the results were normal.

Coronary angiography and left ventriculography were performed in the

following day. The left coronary system was imaged at left and right oblique, right cranial and caudal and anteroposterior cranial positions. The left coronary artery and left ventriculography were normal .The right coronary artery showed significant focal spasm at its proximal segment without provocation that relieved completely by 100 ug intracoronary NTG-.

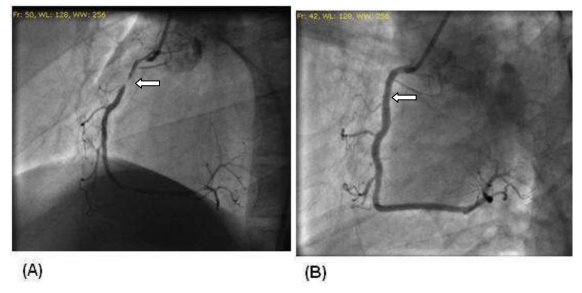


Figure 5. RCA angiogram during spasm (A) and after using NTG (B).

The patient was advised to stop smoking and discharged with LA isosorbide mononitrate 100 mg per day-, amlodipine 5 mg per day-, atorvastatin 20 mg per day. With follow up of more than one year-, the patient feels much better with little symptoms-.

Discussion:

Coronary vasospasm is a transient abnormal contraction of an epicardial coronary artery which can instigate myocardial ischemia.

- 115 Coronary arterial tone varies normally via physiologic mechanisms, but the
- degree of vasoconstriction can range along a spectrum extending from
- undetectable constriction to complete arterial occlusion.
- 118 Many observers use the presence of constriction-induced ischemia as the
- threshold for defining clinical coronary artery vasospasm (3), which has also
- been called vasospastic angina or variant angina.
- It is an important cause of morbidity, but rarely causes mortality.
- 122 Coronary spasm is caused by abnormal coronary smooth muscle activity
- which is not a rare occurrence limited to a particular form of variant
- angina, but a common pathogenic element in ACS (3). It is predominantly
- occurring at rest and usually associated with transient ST-segment elevation
- on the ECG.
- Multiple mechanisms involved as chronic low grade inflammation with
- increased mast cells level (4) and C-reactive protein (CRP) concentrations (5)
- 129 , and endothelial dysfunction (6) that may enhance vascular smooth muscle
- reactivity to agonists as serotonin, histamine and endothelin (7,8).
- Other possible mechanisms include primary vascular smooth muscle cell
- hyperreactivity (9), increase in autonomic nervous activity (10), magnesium
- deficiency (11), and genetic predisposition (12). Nevertheless, the exact
- cellular mechanisms responsible for the spasm remain elusive.
- Before doing coronary angiography, there was high probability that our
- patient has coronary spasm because there was no obvious CAD risk factors
- apart from smoking and mildly increased LDL-C, together with patient's
- atypical chest pain . Moreover, the patient developed ST-segment elevation

- complicated with serious arrhythmia after termination of exercise that takes
- 140 few minutes before completely resolved.
- 141 Unlike atherosclerotic CAD ,patients with variant angina tend to be younger
- in age (13) and chest pain is commonly severe and may be accompanied by
- palpitations or syncope secondary to arrhythmia. As stable angina, vasospasm
- responds by nitrate medication. Serum cardiac troponins may also prove
- unreliable as they may or may not be raised.
- There is no independent predictor of severity of vasospasm and its occurrence.
- 147 It occurs most often from midnight to early morning and is usually not
- induced by exercise in the daytime (14&15). Some studies have shown that
- mild stage exercise is enough to induce variant angina in early hours of the
- morning even multistage exercise fails to do so in the afternoon (3,14) as was
- the case of our patient.
- 152 Is it by nitrate, atropine or by itself coronary spasm was relieved?
- Our patient's hemodynamic decompensation, which developed during the
- exercise recovery phase, was relieved after intravenous administration of
- atropine, a parasympatholytic agent ,that was preceded with using sublingual
- NTG. Patients with coronary artery vasospasm appear to have a heightened
- vasoconstrictor response to acetylcholine as well as an enhanced response to
- the vasodilator effects of nitrates, an observation that is consistent with a
- deficiency of endogenous nitric oxide activity (3).
- During strenuous exercise, sympathetic discharge is maximal, and
- parasympathetic stimulation is withdrawn. In our patient, bradycardia and

- hypotension in the presence of ongoing ischemia due to coronary arterial spasm occurred during the early recovery phase that may resulted from sudden parasympathetic hyperactivity immediately after exercise which could be abolished with atropine.
- Previously, Yasue and colleagues (16) found that pretreatment with intravenous atropine blocked acetylcholine-induced coronary spasms, and they suggested that parasympathetic tone might play a role in the pathogenesis of coronary arterial spasm.
- On the other hand, Wang and associates (17) reported that the isoproterenol head-up tilt test could provoke coronary arterial spasm, and they speculated that both increased basal parasympathetic tone and strong sympathetic stimulation are important in causing coronary arterial spasm.
- Definitive diagnosis is made by angiographically demonstrated coronary 174 artery vasoconstriction either naturally or with provocative tests which 175 reverses with intravenous or intra arterial NTG. In most case reports, the 176 diagnosis was based on the clinical and laboratory findings without 177 provocation (18). A recent guideline by the Japanese Circulation Society 178 Joint Working Group advocated that the diagnosis can be solely established 179 on clinical ground (19). 180 Its management remains a debate with absence of hard scientific evidences 181
- and guidelines. The therapy for vasospastic coronaries can be difficult; up to 25% of patients continue to have intractable angina despite optimal treatment (20). These episodes can be detrimental and occasionally life-threatening when myocardial infarction or arrhythmias occur.

| 186 | Failing medical therapy, mechanical revascularization has been tried |
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| 187 | successfully. Scattered reports of coronary stenting suggest that a |
| 188 | percutaneous strategy may be feasible in such patients (21). |
| 189 | In spite that stent implantation on vasospastic artery bears the danger of in- |
| 190 | stent restenosis and recurrent spasm ,drug-coated stents is favourable as it is |
| 191 | safer and limits the risk of restenosis. The results for surgical revascularization |
| 192 | have been variable, but overall, bypass surgery appears to provide clinical |
| 193 | benefit to less than 50% of patients (22). |
| 194 | In these patients, adding complete plexectomy to the procedure may provide |
| 195 | additional benefit (23). |
| 196 | Mortality though rare, is not uncommon. Long-term survival is believed to be |
| 197 | good, especially in patients who tolerate calcium antagonists and avoid |
| 198 | smoking (24). Predictors of poorer prognosis include the presence of |
| 199 | concurrent coronary atherosclerosis (22),ongoing smoking, intolerance of |
| 200 | calcium antagonists, and spasm of multiple coronary arteries (25). |
| 201 | In conclusion, variant angina can be readily diagnosed by clinical criteria |
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| 202 | and/or provocative testing, yet it is often not considered. Traditionally, such |
| 203 | patients have been reassured that they do not have heart disease despite |
| 204 | persistent symptoms and re-hospitalization. |
| 205 | Given that it can have life-threatening sequelae that are preventable with |
| 206 | readily available therapies, it is essential that clinicians are vigilant in |
| 207 | considering this condition. |
| | |

Abbreviations

- 209 CAD: coronary artery disease; ECG: electrocardiogram; RPP: rate-pressure
- product; CHB: complete heart block; NTG: nitroglycerine; CXR: Chest x-ray;
- BMI: Body mass index; BP: Blood pressure; HR: Heart rate; HDL-C: High
- density lipoprotein-cholesterol; LDL-C: Low density lipoprotein-cholesterol;
- TG: Triglycerides; HBA1c: Glycosylated hemoglobin; NTG: nitroglycerin.

Consent

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- 215 Written informed consent was obtained from the patient for publication of this
- 216 case report.

217 Declaration of Interest

218 The authors report no conflicts of interest

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