CASE REPORT

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Takotsubo cardiomyopathy complicating acute on chronic pancreatitis: A case report



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ABSTRACT

Acute or chronic pancreatitis as a trigger of Takotsubo cardiomyopathy (TCM) has been infrequently described in the literature. Misdiagnosis of this phenomenon can often occur due to overlap in symptomology, particularly in those outside of the usual. A 50-year-old man with a history of alcohol abuse presented with epigastric and chest pain. Electrocardiography showed ischemic changes, and laboratory workup revealed elevated lipase, amylase, and troponin. He was diagnosed with acute pancreatitis and managed presumptively as acute coronary syndrome, class 4 angina. Patient had no signs of chronic liver disease, his pulmomary system was stable. He was a known case of left upper limb thrombosis and had portal venous thrombosis and was on tablet Rivaraxoban 10 mg. Subsequent coronary angiography was negative for obstructive coronary artery disease, and left ventriculography demonstrated basal hyperkinesis and apical akinesis, characteristic of TCM. TCM is a rare complication of acute pancreatitis. Increased awareness of this phenomenon is required to prevent delays in diagnosis and avoid unnecessary interventions and complications.

Key words: Ethanol related acute on chronic pancreatitis; Takotsubo cardiomyopathy

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INTRODUCTION

Takotsubo cardiomyopathy (TCM) has increasingly gained international awareness since it was introduced in 1990. While there are many well-documented triggers of TCM, the role of acute pancreatitis has been only sporadically mentioned in the literature. Increasing awareness of this phenomenon particularly in those outside of the usual patient demographic may lead to earlier diagnosis and avoid unnecessary interventions. We present a rare case of TCM complicating ethanol-related acute on chronic pancreatitis.

CASE PRESENTATION

A 50-year-old man, with a history of smoking and chronic alcoholism and no other cardiac risk factors, developed chest pain with radiation to left shoulder,

epigastric pain associated with nausea, vomiting, and diaphoresis, following significant alcohol consumption the night prior. On presentation, he was tachycardic to 112 beats/min, epigastria tenderness was present. Pain was radiating to back and associated with nausea and vomiting. On examination of his abdomen, there was no distention or guarding and rigidity. His laboratory workup showed an elevated lipase of 221 U/L, leukocytosis of 13,300 Cell/Cu.mm, C-reactive protein of 54 mg/L, and liver functions and renal function was normal. Kidney function was maintained with a creatinine of 0.6 mg%. Chest radiography, when performed, showed features of acute pulmonary edema and electrocardiography (ECG) revealed ST elevation in the anterior leads (Figure 1). A provisional diagnosis of anterior ST-elevation myocardial infarction was made. Echocardiogram (Figure 2) was done showed hypokinesia of left ventricle in the distal 2/3rd of

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interventricularseptum, anterior wall of left ventricle, ejection fraction was 39% with moderate left ventricular systolic function.

Coronary angiography with ventriculography showed nonobstructive coronary artery disease and basal hyperkinesis with apical akinesis of the left ventricle (Figures 3 and 4). The final diagnosis of TCM was made. Dual antiplatelet therapy was ceased.

Contrast-enhanced computed tomography of the abdomen demonstrated an edematous pancreas with calcification, collection, diffuse peripancreatic fat stranding. The patient underwent USG-guided percutaneous drainage of the collection. Therefore, the patient was concurrently diagnosed with complicated acute on chronic pancreatitis with pancreatic fluid collection. He was managed conservatively with intravenous fluids and gradual diet upgrade. The patient's abdominal pain resolved after 2 weeks.

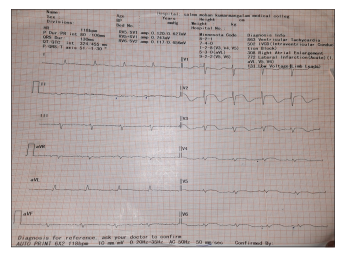


Figure 1: Electrocardiogram



Figure 2: Echocardiography

TCM is characterized by transient left ventricular wall abnormality, resulting in apical¹ ballooning resembling an octopus trap, "takotsubo." Most cases of TCM are associated with a preceding stressor, and¹³ sympathetic stimulation in the setting of increased catecholamines is widely accepted to be central to its pathogenesis. Classically, this association has been with emotional trauma — that is

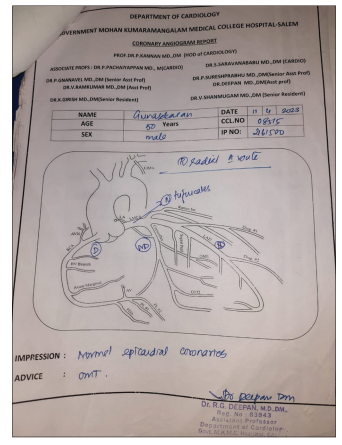


Figure 3: Coronary Angiogram

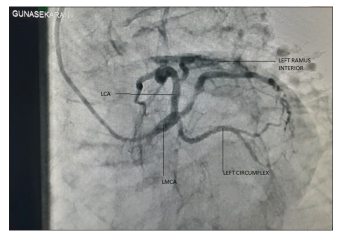


Figure 4: Left oblique anterior view

Image: Second	Table 1: S	Table 1: Summary of literature reporting on pai	erature repo	orting on p	ancreatitis-induced I CIM	-IIIaacea I							
att 56 bit bit bit bit bit bit bit bit bit bit	Authors	Sankri-Tarbichi et al. ²	Rajani et al. ⁷	Cheezum et al.⁴	Pednekar and Chandra ⁶	Leubner et al. ⁸	Bruenjes et al. ¹⁰	Boulos ⁹	Garbowska et al. ¹²	Koop et al. ¹⁴	Abe et al. ¹⁵	Ashraf et al. ¹⁶	Current case
Rudo Abornial N-V RUO pain MeV N-V N-V <	Age (years) Sex Etiology of	56 Female Gallstones	72 Female -	76 Female Gallstones	70 Female -	76 Female Gallstones	55 Male Alcohol	47 Female -	47 Female Alcohol	63 Male Gallstones	57 Female Alcohol	64 Female Unknown	50 Male Alcohol
TCM 3 days 7 days 2 days 2 days 4 days 4 days afion - - WCC - WCC	pancreautis Symptoms of pancreatitis	RUQ pain, nausea	Abdominal pain	>+N	RUQ pain	Epigastric pain, N+V	Epigastric pain, N+V	Epigastric pain		Epigastric pain, N+V	Abdominal pain, N+V	Epigastric pain,	Epigastric pain, N+V
ms of massaa SOB, chrest pain, byposentic Tadrypres, byposentic Candiac arrest approsentic SOB, massaa Tadrypres, SOB, PEA arrest ansea SOB, SOB, PEA arrest ansea SOB, SOB, PEA arrest ansea SOB, and sob ansea SOB, SOB, PEA arrest ansea SOB, and ansea SOB, SOB, PEA arrest ansea S	Time to TCM Systemic inflammation	3 days -	7 days _	2 days WCC 23.3×10º/L	Same day -	1 day Mild leukocytosis	Same day -	1 1	7 days WCC 19.52×10 ⁹ /L,	3 days WCC 11.5×10⁰/L	4 days WCC 14.6×10⁰/L	vomung 5 days -	diaprioresis Same day WCC 13.2×10º/L
(10g 2.39 0.32 0.67 3.13 9.94 0.65 0.3 9.65 0.02 0.02 0.04 renore TWUV2-5 Inferolateral Lateral ST Inferolateral Lateral ST Inferolateral Lateral ST Inferolateral ST Inferolateral ST Inferolateral ST 0.02 0.02 0.02 0.04 Pulmonary - Pulmonary - Pulmonary - Pulmonary Pulmonary phy edema - Pulmonary - - Pulmonary Pulmonary Pulmonary point - Pulmonary - - Pulmonary - Pulmonary Pulmonary point - Pulmonary - - Pulmonary - Pulmonary point - Pulmonary - - Pulmonary Pulmonary Pulmonary point - Pulmonary - - - - Pulmonary Pulmonary point - Pulmonary - - - - Pulmonary Pulmonary point - Pulmonary - - Pulmonary Pulmonary Pulmonary point - Pulmonary	Symptoms of TCM	SOB, chest pain, nausea	Chest pain	Tachypnea, hypoxemic	Cardiac arrest	SOB, diaphoresis	Chest pain, diaphoresis,	Nausea	CKP 293.8 mg/L Chest pain, SOB	Oliguria, hypotension, sood DEA accord	SOB, hypoxemic	SOB	Chest pain
TWI V2-5 Inferolateral TW Lateral ST Inferior ST Anteroseptal Generatised anterior TW Inferolateral ST Interolateral ST	Troponin (ng/ mL; reference	2.39	0.32	0.67	3.13	9.94	nausea 0.66	0.3	9.65	o.02 0.02	0.97	Elevated	1019.63
est Pulmonary cedema codema pulmonary cumonary c	< U.UZ) ECG	TWI V2-5	Inferolateral TWI	Lateral ST elevation	Inferior ST elevation, anterior TWI	Anteroseptal ST elevation		Inferolateral TWI	ST elevation V2	Non-specific inferolateral T-wave	Diffuse ischemic TWI	Anterior ST elevation	Anterior ST elevation
hocardiogramLVEF 25%, ancalApical akinesisLVEF 30%, akinesisLVEF 30%, anterior, basalLVEF 20-25%, apicalLVEF 20-25%, apical<	Chest radiography	Pulmonary oedema	I	Pulmonary oedema, bilateral pleural effusions	1	Mild pulmonary oedema, bilateral pleural	I	I	Pulmonary congestion	crianges Acute pulmonary oedema	Pulmonary oedema	Pulmonary oedema	I
Dasse Dasse Ventrice Normal coronary Unobstructed Mild non- No obstructive No obstructive Normal coronary S0% LAD Normal arteries coronary obstructive atherosclerotic CAD Not done- Normal coronary S0% LAD Normal arteries coronary obstructive atherosclerotic CAD myocardial atteries coronary arteries CAD disease coronary obstructive stenosis, coronary Aspirin, BB, ACEi BB, ACEi - Aspirin, BB, - - Left ventricular BB, ACEi Yes - Yes - - - Left ventricular BB, ACEi Yes - Yes - - - Left ventricular BB, ACEi 10 days - 2 weeks - 3 weeks - 10 days	Echocardiograr or ventriculograph	n LVEF 25%, severe apical y hypokinesia/ akinesia of left ventricle, hypercontractile	Apical akinesis	LVEF 30%, severe apical hypokinesis + hyperdynamic basal contraction		effusions LVEF 30–35%, hypokinetic apical left ventricle	LVEF 25%, apical ballooning, hypercontractile basal segments	Akinesis of distal anterior, lateral, inferior walls of left	LVEF 25%, apical ballooning, hypercontractile basal segments of left ventricle	LVEF 20–25%, new-onset cardiomyopathy, global hypokinesis	LVEF 40%, basal segment hyperkinesis, apical akinesis		LVEF 39% hypokinesia of left ventricle
nt of Aspirin, BB, ACEi BB, ACEi BB, ACEi - Aspirin, BB, Left ventricular BB, ACEi variarin v fes - Yes Yes Yes No 10 days - 2 weeks 6 weeks - 3 weeks - 10 days 3 weeks -	Angiography	base Normal coronary arteries	Unobstructed coronary arteries	U	No obstructive atherosclerotic disease	No CAD	No obstructive CAD	ventricle Not done— myocardial nuclear stress test mildly	Normal coronary arteries	50% LAD stenosis, otherwise no obstructive CAD	Normal coronary arteries	to akinetic Only luminal irregularities	Normal epicardial arteries
10 days – 2 weeks 6 weeks – 3 weeks – 10 days 3 weeks –	Treatment of TCM Recovery of	Aspirin, BB, ACEi Yes		BB, ACEi Yes	BB, ACEi Yes	1 1	Aspirin, BB, ACEi, warfarin Yes		- Yes	Left ventricular assist device Yes	BB, ACEi No	BB, ACEi Yes	BB, ACEi -
recovery	LVEF Time to recovery	10 days	I	2 weeks	6 weeks	I	3 weeks	I	10 days	3 weeks	I	6 weeks	I

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the "broken heart"; however, physical stressors are in fact more common. These include various physical activities, procedures, drugs, and medical conditions ranging from sepsis to lightning strike. Modified Mayo Criteria is used to identify TCM. Other criteria used are Japanese guidelines 2007, Gothenbeurg criteria – Sweden 2012, John Hopkins criteria 2012, Madias criteria 2014.

Acute pancreatitis as a trigger of TCM is a rare phenomenon. Since it was first described in 2007, there have been only 11 total cases reported in the literature (Table 1). Of these, 9 were female, with ages ranging between 47 and 76 years and a median of 63 years, and 82% of cases above 50 years. The most common etiology for acute pancreatitis is gall stones and alcohol consumption. The etiology of pancreatitis reflects that typically seen in the community with alcohol and gallstones predominating. The timeframe for onset of pancreatitis symptoms to TCM was quite variable, ranging from hours to 7 days. Similarly, the markers of systemic inflammation were an unreliable predictor of the development of TCM or its complications, with leukocytosis ranging from mild to markedly elevated.

Recognizing TCM in the setting of pancreatitis is clinically difficult, TCM complicating pancreatitis results in considerable overlap in symptomology, and a patient's epigastric pain can easily mask or distract from chest pain. Formal diagnosis of TCM relies heavily on coronary angiography with left ventriculography.^{3,11} While multiple diagnostic criteria have been proposed, the most widely known is the Revised Mayo Clinic Criteria.¹¹ This requires the presence of transient left ventricular dysfunction, the absence of obstructive coronary artery disease, electrocardiographic abnormalities, or troponin elevation.

All previously reported cases of pancreatitis-induced TCM, and our own case, had elevated troponin, and ECG almost always demonstrated ST elevation or T-wave inversion. Chest radiography, when performed, showed features of acute pulmonary edema. Furthermore, almost all cases demonstrated left ventricular apical hypokinesis or akinesis, basal hyperkinesis, reduced ejection fraction, and normal or non-obstructive coronary arteries.¹⁴

Diagnostic challenge lies in the overlap of biochemical and ECG changes of TCM with those of acute coronary syndrome. Our presented case, like several others, was diagnosed presumptively as acute coronary syndrome. The patient underwent coronary angiogram which showed normal epicardial arteries, ruling out acute coronary syndrome.

However, in a center where primary percutaneous coronary intervention was available within 90 min, a diagnosis of TCM could have been reached sooner, and the potential complications of thrombolysis could have been avoided. Interestingly, there may be a role in the stable patient for early echocardiogram and subsequent computed tomography coronary angiography to reach a diagnosis of TCM, thereby avoiding the vascular complications of cardiac catheterization.

CONCLUSION

The outcomes of TCM as a complication of pancreatitis appear to be good. There have been no reported deaths, even in the severe case of cardiac arrest ultimately requiring left ventricular assist device.

All patients, where treatments were described, were discharged on a beta-blocker and ACEi, and in those whose follow-up was reported, all but one had recovered left ventricular function, ranging from as early as 10 days up to 6 weeks. Abe et al.,¹⁵ reported the case of persistently reduced left ventricular function even at 4 months, which was hypothesized to be due to ongoing alcohol use.

Overall, TCM is a rare, and potentially under-recognized, complication of acute pancreatitis. In a scenario where delays in diagnosis can lead to unnecessary interventions and complications, increasing awareness of this phenomenon by surgeons and physicians alike is imperative.

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