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Takotsubo syndrome (TTS) is a poorly recognized heart disease that was initially regarded as a benign condition. Recently, it has been shown that TTS may be associated with severe clinical complications including death and that its prevalence is probably underestimated. Since current guidelines on TTS are lacking, it appears timely and important to provide an expert consensus statement on TTS. The clinical expert consensus document part I summarizes the current state of knowledge on clinical presentation and characteristics of TTS and agrees on controversies surrounding TTS such as nomenclature, different TTS types, role of coronary artery disease, and etiology. This consensus also proposes new diagnostic criteria based on current knowledge to improve diagnostic accuracy.

Keywords
- Takotsubo syndrome
- Broken heart syndrome
- Takotsubo definition
- Acute heart failure
- Consensus statement
- InterTAK Diagnostic Criteria

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History

The term takotsubo syndrome (TTS) was first introduced when Sato et al. published their report of five cases in a Japanese textbook in 1990. The first TTS case of this series was managed in 1983 in the Hiroshima City Hospital (Figure 1). A 64-year-old female presented with acute chest pain consistent with acute myocardial infarction (AMI), typical electrocardiographic (ECG) changes, but normal coronary arteries and an unusual appearance of the left ventricle (LV) with a narrow neck and apical ballooning during systole. Interestingly, the marked wall motion abnormalities on left ventriculography disappeared after 2 weeks. Over time TTS was more frequently diagnosed in Japan. Therefore, it was first assumed that this disorder only affected people of Asian descent, as TTS was completely unknown to the Western world until the first cases were published from French and American research groups in the late 1990s. Desmet et al. introduced the first patient case series in Caucasians using the term ‘takotsubo’.

Takotsubo syndrome gained international awareness among researchers and physicians when Wittstein et al. reported their findings in the New England Journal of Medicine in 2005. Since then TTS has been more frequently recognized worldwide but still remains an underappreciated and often misdiagnosed disorder.

Nomenclature

Takotsubo syndrome derived its name from the Japanese word for octopus trap, due to the shape of the LV at the end of systole and has been described under a remarkable number of different names in the literature including 'broken heart syndrome', 'stress cardiomyopathy', and 'apical ballooning syndrome'. No single term precisely describes the heterogeneous ventricular appearance with which this syndrome can occur. To date, consensus has not been reached on the nomenclature. The term ‘takotsubo’ is widely used in acknowledgement of the
Japanese physicians who initially described this disorder. However, in contrast to other cardiomyopathies that are usually not transient in nature, TTS is characterized by a temporary wall motion abnormality of the LV and shares common features with acute coronary syndrome (ACS) [similar symptoms at presentation, ECG abnormalities, elevated cardiac biomarkers as well as a comparable in-hospital mortality with ST-segment elevation myocardial infarction (STEMI) and non-STEMI] specifically in terms of a microvascular ACS form. Among different etiologies of heart failure such as coronary artery disease (CAD), tachyarrhythmias etc. TTS includes a wide spectrum of emotional or physical triggers resulting also in left ventricular dysfunction. Therefore, it is best described as a ‘syndrome’ and the term ‘takotsubo syndrome’ seems most appropriate. 

**Epidemiology**

Since the initial report by Japanese cardiologists 25 years ago, TTS has been increasingly recognized in diverse countries across six continents. Takotsubo syndrome is estimated to represent approximately 1–3% of all and 5–6% of female patients presenting with suspected STEMI. The Nationwide Inpatient Sample discharge records from 2008 using the International Classification of Diseases revealed that TTS accounts for 0.02% of hospitalizations in the United States. Recurrence rate of TTS is estimated to be 1.8% per-patient-year. Based on the published literature about 90% of TTS patients are women with a mean age of 67–70 years, and around 80% are older than 50 years (Figure 2). Women older than 55 years have a five-fold greater risk of developing TTS than women younger than 55 years and a 10-fold greater risk than men. With growing awareness of TTS, male patients are diagnosed more often, especially after a physical triggering event. TTS has also been described in children with the youngest reported TTS patient being a premature neonate born in the 28th gestational week. Current data on racial differences are inconsistent and large-scale studies are lacking. However, it has been reported that TTS seems to be uncommon in African–Americans and Hispanics, while most of the cases reported in the United States have been Caucasians. Furthermore, it has been reported that patients of African-American descent have more in-hospital complications such as respiratory failure, stroke and require more frequently mechanical ventilation compared to Caucasians and Hispanics. With regard to ECG differences, it has been shown that QT prolongation as well as T-wave inversion are more often reported in African-American women with TTS. Of note, regarding gender differences the TTS prevalence in men appears to be higher in Japan. The prevalence of TTS appears to be higher in patients with non-emotional triggers admitted to intensive care units. Moreover, it is likely that
Symptoms and signs

The most common symptoms of TTS are acute chest pain, dyspnoea, or syncope and thus indistinguishable from AMI at the first glance. However, in some patients, TTS may be diagnosed incidentally by new ECG changes or a sudden elevation of cardiac biomarkers. Clinical manifestation of TTS induced by severe physical stress may be dominated by the manifestation of the underlying acute illness. In this regard, patients with ischaemic stroke or seizure-triggered, TTS had less frequent chest pain, which could be explained by impaired consciousness, neurologic complications, or a sudden haemodynamic deterioration. In contrast, patients with emotional stress factors had a higher prevalence of chest pain and palpitations. Importantly, a subset of TTS patients may present with symptoms arising from its complications, e.g. heart failure, pulmonary oedema, stroke, cardiogenic shock, or cardiac arrest.

Diagnostic criteria

The diagnosis of TTS is often challenging because its clinical phenotype may closely resemble AMI regarding ECG abnormalities and biomarkers. While a widely established non-invasive tool allowing a rapid and reliable diagnosis of TTS is currently lacking, coronary angiography with left ventriculography is considered the gold standard diagnostic tool to exclude or confirm TTS. Abe et al. introduced the first diagnostic criteria for TTS in 2003. One year later, a dedicated group of cardiologists from the Mayo Clinic proposed their diagnostic criteria. In 2006, the American College of Cardiology and American Heart Association classified TTS as a primary acquired cardiomyopathy. In 2008, the revised version of the Mayo Clinic Diagnostic Criteria was published incorporating neurogenic stunned myocardium. Furthermore, the authors defined different TTS sub-types and highlighted that obstructive coronary lesions may occasionally be present concomitantly. The Mayo Clinic Diagnostic Criteria are the most widely known, but exceptions to the rule [e.g. the presence of CAD] are poorly appreciated among physicians and cardiologists. More recently, other research groups have proposed slightly different criteria for TTS, i.e. the Japanese Guidelines, the Gothenburg criteria, the Johns Hopkins criteria, the Tako-tsubo Italian Network proposal, the criteria of the Heart Failure Association (HFA) TTS Taskforce of the European Society of Cardiology (ESC) as well as the criteria recommended by Madias. Thus, there is a lack of a worldwide consensus. Based on current knowledge, we have developed new international diagnostic criteria (InterTAK Diagnostic Criteria, Table 1) for the diagnosis of TTS that may help to improve identification and stratification of TTS. The most important changes with accompanying rationale include:

(i) Pheochromocytoma is a neuroendocrine tumour derived from enterochromaffin cells of the adrenal gland that may lead to a ‘catecholamine storm’ with LV dysfunction, ECG abnormalities, and increased biomarkers as well as hypercontraction of sarcomeres and contraction band necrosis indistinguishable from TTS. Pheochromocytoma is also included as a secondary cause of TTS in the diagnostic criteria of the HFA of the ESC.

Notwithstanding, most of the diagnostic criteria have excluded pheochromocytoma as a specific cause of TTS. The Japanese criteria emphasize that pheochromocytoma is a TTS-like myocardial dysfunction. Pheochromocytoma is also included as a secondary cause of TTS in the diagnostic criteria of the HFA of the ESC.

(ii) Concomitant CAD is reported with a prevalence ranging from 10–29%. In this regard, patients with TTS and obstructive CAD are often misdiagnosed as classical ACS and differentiation can be challenging. Therefore, the presence of CAD should not be considered as an exclusion criterion as acknowledged by the modified...
Pathophysiology

Sympathetic stimulation

The precise pathophysiological mechanisms of TTS are incompletely understood, but there is considerable evidence that sympathetic stimulation is central to its pathogenesis. An identifiable emotionally or physically triggering event precipitates the syndrome in most cases.66 and TTS has been associated with conditions of catecholamine excess (e.g. pheochromocytoma),64 central nervous system disorders,64 and activated specific cerebral regions.65 Clinical features of TTS and the various ballooning patterns can be caused by intravenous administration of catecholamines and beta-agonists.66 Although it has been shown that patients with TTS triggered by emotional stress have markedly elevated levels of catecholamines compared to patients with Killip Class III myocardial infarction,67 others68 could not replicate this finding most likely due to methodological issues. In line with a sympathetic stimulation, elevated norepinephrine levels in the coronary sinus have been found in TTS patients, suggesting an increase in the local release of myocardial catecholamines.69 Accordingly, analyses of heart rate variability have also demonstrated a sympathetic predominance and marked depression of parasympathetic activity during the acute phase.70 Microneurographic studies confirmed increased muscle sympathetic nerve activity and decreased spontaneous baroreflex control of sympathetic tone in some TTS patients.60 as did myocardial scintigraphy using 123I-metaiodobenzylguanidine.67 Furthermore, abnormalities in myocardial sympathetic function can persist for months after recovery of LV systolic function.62 These abnormalities appear to induce an interstitial mononuclear inflammatory response and occasionally contraction band necrosis.5

Several animal models have also supported the central role of adrenergic stimulation in TTS.63–66 In rats, LV apical ballooning can be provoked by immobilization stress and attenuated by alpha- and beta-receptor blockade.68 Furthermore, in a more recent and novel rat model, it was possible to demonstrate that the administration of different catecholamines instigates the various ventricular ballooning patterns by an afterload-dependent mechanism.67

Potential pathophysiological effects of enhanced sympathetic stimulation

Although enhanced sympathetic stimulation is central to TTS, the mechanism by which catecholamine excess precipitates myocardial stunning in the variety of regional ballooning patterns that characterize this syndrome is unknown. Several hypotheses have been proposed as follows:

Plaque rupture

It has been suggested that transient ischaemia induced by plaque rupture followed by rapid lysis may cause myocardial stunning in patients with apparent non-obstructed CAD at angiography. Indeed, eccentric atherosclerotic plaques in the mid-portion of the left anterior descending (LAD) coronary artery have been reported, but intravascular ultrasound and optical coherence tomography have failed to identify ruptured plaques in the vast majority of TTS patients.68–70 Furthermore, this explanation is very unlikely as patients with TTS exhibit wall motion abnormalities extending beyond single coronary vascular territories and also sometimes include the right ventricle. In addition, the apical ballooning phenotype is known to occur in the absence of a wraparound LAD and this coronary anatomical variant is not more prevalent in TTS than in the control group.71

Multi-vessel epicardial spasm

Sympathetically mediated epicardial spasm has been proposed as a potential cause in TTS. Takotsubo syndrome may be associated with endothelial dysfunction and other conditions of abnormal vasomotor function such as migraine or Raynaud’s phenomenon.72 Similarly, endothelium-dependent dilation is reduced after emotional stress and prevented by endothelin antagonists.73 At presentation, patients with TTS have marked impairment in brachial artery flow-mediated dilation compared to those with infarction or healthy controls, which gradually improves over several weeks.74 In the early recovery period, predisposition to coronary vasospasm using intracoronary acetylcholine was demonstrated in some, but not all TTS patients.75 Furthermore, it has been suggested that the pattern of LV dysfunction in patients with TTS may require involvement of specific coronary side branches.76 Similarly, myocardial bridging in the LAD has been considered.77 Although epicardial coronary vasoconstriction may contribute to TTS in a subset of patients,78 the vast majority of patients do not show any evidence of epicardial spasm even with use of provocative agents.

Furthermore, endothelial dysfunction is often associated with oxidative stress, and studies suggest that this may play a role in myocardial dysfunction in TTS. A recent study by Zhang et al.79 found that hydrogen sulfide relieved cardiac dysfunction in animal models by decreasing oxidative stress. It has been reported that the level of oxidative stress correlates to the extent of myocardial dysfunction in TTS patients in the acute recovery phase. Nanno et al.80 measured 8-hydroxy-2’-deoxyguanosine (8-OHdG) and norepinephrine levels in TTS patients compared with AMI patients. They found that 8-OHdG levels changed proportionately with wall motion score and plasma levels of norepinephrine were twice as high in TTS patients as in AMI patients.

Microcirculatory dysfunction

Catecholamines and endothelin exert their vasoconstrictor effects primarily in the coronary microvasculature where 1-receptors81 and endothelin receptor type A predominate suggesting that acute microcirculatory dysfunction may have a central role in TTS.
Furthermore, acutely TTS exhibits decreased microRNA (miRNA) 125a-5p as well as increased plasma levels of its target endothelin-1 in line with the microvascular spasms hypothesis. Microvascular blood flow may be reduced in the acute phase of TTS as is coronary flow reserve. Similarly, increased thrombosis in myocardial infarction (TIMI) frame counts and abnormal grades of TIMI myocardial perfusion have been noted.

In the acute phase, intravenous administration of adenosine has been shown to transiently improve myocardial perfusion, wall motion score index, and left ventricular ejection fraction (LVEF) in TTS, suggesting that intense microvascular constriction plays a major role in the pathophysiology. In addition, the notion of acute microcirculatory dysfunction in TTS as a contributing pathophysiological factor secondary to enhanced sympathetic stimulation is supported by endomyocardial biopsies revealing apoptosis of microvascular endothelial cells. Microcirculatory dysfunction in the acute phase of TTS is transient and its recovery appears to correlate with improved myocardial function.

Cold pressor testing 1–3 years after the acute episode results in an elevation of catecholamines and transient apical and mid-LV wall motion abnormalities. Mental stress or reactive hyperaemia result in lower vasomotor responses, but higher catecholamine levels in women with TTS compatible with impaired vascular reactivity and endothelial function. Similarly, in women with a history of TTS coronary vasomotion to acetylcholine is impaired. Impaired microvascular endothelial function was observed in virtually all patients with TTS.

**Catecholamine toxicity on cardiomyocytes**

Transient LV dysfunction in TTS could also result from direct effects of catecholamines on cardiomyocytes. Endomyocardial biopsies revealed occasional contraction band necrosis, which is generally observed in clinical settings of extreme catecholamine production such as pheochromocytoma or subarachnoid haemorrhage, associated with hypercontracted sarcomeres, dense eosinophilic transverse bands, and interstitial mononuclear inflammation as a reflection of myocyte injury. Catecholamines can decrease myocyte viability through cyclic adenosine monophosphate (cAMP) mediated Ca\(^{2+}\) overload as it may occur in TTS. Sarcoplasmic-Ca\(^{2+}\) -adenosine-triphosphatase (SERCA2a) gene expression is downregulated and that of sarcoplin upregulated, while phospholamban is dephosphorylated in TTS. Thus, an increased phospholamban/SERCA2a ratio could result in contractile dysfunction due to decreased Ca\(^{2+}\)-affinity. Indeed, intense G-protein stimulated β-adrenergic receptor signalling modulates gene expression via the cAMP responsive element binding protein-1 and nuclear factor of activated T-cells signalling pathways.

In rodent heart failure models, administration of isoproterenol yields apical fibrosis, and abnormalities of apical contraction and metabolism, features known to occur in dysfunctional apical segments during the acute phase in TTS using fludeoxyglucose-positron emission tomographic studies. In animal models, intracellular lipid droplets accumulate in cardiomyocytes in response to high doses of catecholamines, as in endomyocardial biopsies of TTS patients during the acute phase, but not after recovery. In a rat model of TTS myocardial perfusion in dysfunctional segments appears preserved, challenging microvascular spasm as a primary mediator.

In the mammalian LV β-adrenergic receptor density is highest in the apex, while sympathetic innervation is the lowest suggesting that it may be more sensitive to high levels of catecholamines which may reduce not only coronary blood flow, but at high levels paradoxically also exert negative inotropic effects due to a ‘molecular switch’ of the β-adrenergic receptor from the positive inotropic G\(_i\) to the negatively inotropic G\(_s\) pathway. Since the β\(_2\)-adrenoreceptor is linked via G\(_i\) activation to stimulation of endothelial nitric oxide (NO) synthase, it seems possible, that peroxynitrate mediated nitrosative stress could lead to negative inotropy and inflammation in TTS. Indeed, TTS patients have markers of increased NO signalling and post-mortem hearts of TTS patients also demonstrate markers of increased nitrosative stress. Peroxynitrite release would also result in activation of poly(ADP-ribose)-transferase-1, which might contribute to the myocardial energetic impairment, which has recently been reported in patients with TTS. Endomyocardial biopsies in patients with TTS further suggest that these anti-apoptotic pathways are activated acutely. A polymorphism of the G-protein receptor kinase 5 (GRK5) gene L41Q that blunts β\(_2\)-G\(_i\) trafficking appears common in TTS. On the other hand, a larger study failed to support the conclusions of this study.

In summary, current evidence suggests that TTS is caused by an acute release of catecholamines from either sympathetic nerves, the adrenal medulla, or as drug therapy and occurs primarily in subjects with increased susceptibility of the coronary microcirculation and of cardiac myocytes to the stress hormones leading to prolonged but transient LV dysfunction with secondary myocardial inflammation.

**Activation of myocardial survival pathways**

The severe wall motion abnormalities seen in TTS are transient suggesting that protective mechanisms are likely to operate to preserve myocardial integrity. Two different mechanisms might elicit myocardial protection. The first one is represented by adrenoreceptor-related protective mechanisms. Indeed, supra-physiological levels of epinephrine trigger β\(_2\)-adrenoreceptor to switch from G\(_s\) to G\(_i\) coupling, thus causing a negative inotropic response, which limits the degree of acute myocardial injury in response to the catecholamine surge. The second mechanism is represented by the phosphoinositide 3-kinase/protein kinase B (AKT) survival pathway, which has been found to be transiently activated during the acute phase of TTS. AKT is critical for postnatal cardiac growth and coronary angiogenesis. Also, its downstream targets, especially the mechanistic target of rapamycin and glycogen synthase kinase 3 (GSK3), are well-established regulators of metabolism, proliferation, and cell survival. Cell survival is achieved through various mechanisms: (i) direct inhibition of apoptosis, (ii) inhibition of proapoptotic transcriptional factors, (iii) enhancement of anti-apoptotic transcriptional factors, and (iv) enhancement of cell metabolism by inhibition of the GSK3.

The demonstration that down-regulation of myocardial function is a protective mechanism caused by a severe reduction of perfusion is confirmed by several clinical studies showing ‘inverse perfusion-metabolism mismatch,’ which is typically observed during myocardial stunning.
Predisposition and risk factors

Psychological and physical stressors are universal and affect virtually all individuals throughout their life. However, very few people develop TTS and even fewer experience recurrent episodes. These observations support the existence of risk factors that may make certain individuals more susceptible to TTS. Predisposition and risk factors for TTS are reviewed below:

Hormonal factors

The striking preponderance of postmenopausal females suggests a hormonal influence. Potentially, declining oestrogen levels after menopause increase the susceptibility to TTS in women. Indeed, women older than 55 years have an almost five-fold risk of developing TTS compared to those younger than 55 years. Oestrogens can influence vasomotor tone via up-regulation of endothelial NO synthase. Also, there is evidence that oestrogens can attenuate catecholamine-mediated vasoconstriction and decrease the sympathetic response to mental stress in perimenopausal women. In women with subarachnoid haemorrhage, low levels of oestriol have been associated with an increased risk of LV wall motion abnormalities. In ovariectomized rats subjected to immobilization stress, ECG and contractile abnormalities can be induced and attenuated with oestrogen supplementation. However, systematic data demonstrating a clear link between oestrogen levels and the development of TTS are lacking so far.

Genetic factors

A genetic predisposition to TTS has been suggested by a report of five cases of familial TTS, two in mother-daughter pairs and three in pairs of sisters. Takotsubo syndrome does not appear to have a multigenerational Mendelian inheritance pattern. Hence, it is likely that a genetic predisposition (if present) may interact with environmental factors, polygenic aetiology and/or recessive susceptibility alleles. Polymorphisms in adrenergic genes indeed affect receptor function and downstream signalling, and this raises the possibility that their distribution may differ in TTS patients. Indeed, functional variants of adrenergic receptor genes have been associated with the magnitude of cardiac dysfunction in patients with subarachnoid haemorrhage and pheochromocytoma, conditions which can trigger TTS.

Beta-1-adrenergic receptor (amino acid position 389) and beta-2-adrenergic receptor (amino acid position 27) variants were associated with a greater release of troponin I and alpha-2-adrenergic receptor deletion (del322–325) with reduced LVEF. However, alpha-2C-adrenergic receptor and beta-1-adrenergic receptor polymorphisms do not seem to differ between TTS and controls. In contrast, a different distribution of beta-1-receptor polymorphisms Arg389Gly [homozygous arginine (Arg)/Arg] is more frequently found in TTS, while beta-2-receptor polymorphisms Gln27Glu [homozygous glutamine (Gln)/Gln] were found more frequently in healthy controls, and no difference was observed in the beta-2-receptor Arg16Gly variant between groups. Furthermore, similar genetic polymorphisms in the beta-1-adrenergic receptor and the beta-2-adrenergic receptor were noted in TTS and controls, while a higher frequency of rs17098707 polymorphism in the GRK5 gene was found in TTS patients. Unfortunately, these studies provide conflicting results and are limited in their gene-targeted approach and incomplete in genetic characterization of the complex adrenergic signalling network. Whole-exome sequencing in 28 TTS subjects revealed no difference in allele frequency or burden between TTS subjects and population controls. As such, these data do not provide strong evidence for a genetic predisposition in TTS, but lend support to genetic heterogeneity and a potential polygenic susceptibility conferring a cumulative effect on dysregulation of adrenergic pathways. Most of the published studies were conducted in small cohorts and much larger cohorts are required to evaluate the genetics of TTS comprehensively.

Borchert et al. have investigated a genetic predisposition for TTS by creating the first ‘takotsubo in a dish’ model by using TTS-specific induced pluripotent stem cell-derived cardiomyocytes (iPSC-CMs). This model recapitulates some of the pathophysiology observed in patients during the acute phase of TTS allowing further exploration of underlying mechanisms. They found an overactive beta-adrenergic pathway and higher sensitivity of catecholamines in TTS compared to population controls. Interestingly, receptor desensitization and different beta1/beta2-adrenoreceptor responses shed further light on the mechanisms of TTS. Based on this TTS-model future treatment targets should be identified to rescue patients with TTS.

Psychiatric and neurologic disorders

A high prevalence of psychiatric and neurologic disorders has been reported in patients with TTS. In an age- and sex-matched comparison between patients with TTS and ACS, rates of psychiatric or neurologic disorders were substantially higher in TTS. Indeed, 27% had an acute, former, or chronic history of neurologic disorders and 42% had a psychiatric diagnosis with half of them suffering from depression. Indeed, anxiety and depression appear more common in TTS than in patients with STEMI or healthy controls in a prospective study, the prevalence of depression and anxiety was 78%, much higher than in patients with ACS. Patients with TTS also appear to have a high prevalence of type-D-personality, which is characterized by negative emotions and social inhibition, and which has been associated with an increased cardiovascular risk. However, another study found no difference in the personality profile and stress coping skills between TTS patients and population controls.

Interestingly, in a recent study comparing the signature of circulating miRNAs in TTS and STEMI, miR-16 and miR-26a, known to be associated with neuropsychiatric conditions, were significantly upregulated in TTS. Psychological disorders may thus have a pathogenic role. Of note, depressed patients have an exaggerated norepinephrine response to emotional stress, and a subset of patients has an increased spillover and decreased reuptake of norepinephrine. Similarly, patients with panic disorder and anxiety have a decreased catecholamine reuptake due to impairment of norepinephrine reuptake transporters. On the other hand, antidepressants, e.g. selective norepinephrine reuptake inhibitors, may facilitate myocardial stunning by increasing local levels of catecholamines. This increased sympathetic response to acute stress combined with greater cardiac sympathetic sensitivity may make patients with mood disorders and anxiety susceptible to stress-related cardiac dysfunction. Takotsubo syndrome has been reported to occur after neurologic disorders especially stroke, subarachnoid haemorrhage, and seizures. Histopathological findings of autopsied patients with sudden unexpected death in epilepsy revealed contraction band...
Triggers

A hallmark of TTS is its association with a preceding stressful event. Initially, most reported triggers involved an emotional trauma. As TTS became more known, an association with physical stressors was also noted as well as TTS cases that occur in the absence of an evident stressor. A systematic illustration of preceding emotional and physical stressors is shown in Figure 3.

Physical triggers are more common than emotional stress factors. Interestingly, male patients are more often affected from a physical stressful event, while in women an emotional trigger can be more frequently observed. Of note, precipitating triggers may represent a combination of emotional and physical issues (e.g., panic attack during a medical procedure), as well as environmental triggers such as long-term exposure to aircraft noise. On the other hand, about one-third of patients presents without evidence of an identifiable preceding stressful event.

In hospitalized patients, TTS may have an atypical presentation and manifest itself by tachycardia, hypotension, heart failure, elevation of enzymes, and physical abnormalities as shown in Figure 3.165

Types of takotsubo syndrome

Although several anatomical TTS variants have been described four major types can be differentiated based on the distribution of regional wall motion abnormalities as shown in Figure 4.16,51 The most common TTS type and widely recognized form is the (i) apical ballooning type also known as the typical TTS form, which occurs in the majority of cases. Over the past years, atypical TTS types have been increasingly recognized. These include the (ii) midventricular, (iii) basal, and (iv) focal wall motion patterns. Recently, it has been demonstrated that patients suffering from atypical TTS have a different clinical phenotype. These patients are younger, suffer more often from neurologic comorbidities, have lower brain natriuretic peptide values, a less impaired LVEF, and more frequent ST-segment depression compared to typical TTS. In-hospital complication rate is higher in typical TTS. After adjustment for confounders, LVEF <45%, atrial fibrillation, neurologic disorders but not TTS phenotype were independent predictors of death. Beyond 1-year, long-term mortality is similar in typical and atypical TTS phenotypes, therefore, patients should be equally monitored and treated. The basal phenotype has been reported to be associated with the presence of phochromocytoma, epinephrine-induced TTS, or subarachnoid haemorrhage consequently, these conditions should be considered in this particular setting.

Besides the four major TTS types, other morphological variants have been described including the biventricular (apical type and right
ventricular involvement),\textsuperscript{19} isolated right ventricular,\textsuperscript{188,189} and global form.\textsuperscript{190} Global hypokinesia as a manifestation of TTS is difficult to prove given the very broad differential diagnoses including conditions such as tachycardia-induced cardiomyopathy. Right ventricular involvement is present in about one-third of TTS patients and may be a predictor for worse outcome.\textsuperscript{191} The true prevalence of the isolated right ventricular form is unknown since little attention is paid to the right ventricle in daily clinical echo routine.

Figure 3 Emotional and physical stress factors precipitating takotsubo syndrome. Reprinted, modified, and translated with permission from Schlossbauer et al.\textsuperscript{7} COPD, chronic obstructive pulmonary disease; PRES, posterior reversible encephalopathy syndrome; TIA, transient ischaemic attack.
Figure 4. The four different types of takotsubo syndrome during diastole (left column) and systole (middle column). The right column depicts diastole in red and systole in white. The blue dashed lines demonstrate the region of the wall motion abnormality. Reprinted and modified with permission from Templin et al.\textsuperscript{16}
Patients with recurrent TTS can demonstrate different wall motion patterns at each event, suggesting that left ventricular adrenergic receptor distribution does not explain different TTS types.

Chronobiology

A growing body of evidence reveals that acute cardiovascular events are not distributed randomly over time, but instead depend on the time of day, day of the week, and months/seasons of the year. Several studies have investigated chronobiological features of TTS. Two studies reported a peak in the morning and afternoon, while others failed to show a statistically significant temporal pattern. Two studies observed the highest frequency on Monday and a third investigation has not found a weekly variation. Most conducted studies reported a summer preference for TTS, while one study reported a winter peak. Hence, conflicting results about the chronobiological pattern of TTS exist.

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