Echocardiographic Findings in Heart Failure Patients With Methamphetamine Use: A Case-Control Study

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Abstract

Background
Methamphetamine use is associated with cardiovascular disease and significant morbidity and mortality. There is only one previous study performed on echocardiographic parameters in patients with methamphetamine cardiomyopathy.

Methods
We performed a retrospective review of medical records in a county hospital in Southern California with a high population of methamphetamine users. We reviewed medical records and echocardiogram findings in patients seen in our institution from November 2019 to November 2020 who had cardiomyopathy with and without methamphetamine use. We excluded patients who either left the hospital or expired before appropriate assessment. We divided our patient population into a case group (methamphetamine users) and a control group (non-methamphetamine users) to study and compare their echocardiographic parameters.

Results
Case group included a total of 254 patients and control group included 268 patients. Majority of the patient population were males - 178 (70%) and 180 (67%) in the case and control group respectively. Age was found to be statistically significant with the younger population in the case group (p = 0.0000). Our analysis revealed statistically significant difference in methamphetamine users compared to non-users in regards to left ventricle ejection fraction (33.65% ± 18.02 vs. 41.55% ± 15.61, p=0.0000), left ventricle mass index (122.49 grams/m² ± 40.66 vs. 108.62 grams/m² ± 32.82, p=0.0000), left ventricle end diastolic volume index (85.91 mL/m² ± 37.40 vs. 72.44 mL/m² ± 25.44; p=0.0000) and marginally significant right ventricle systolic pressure (42.29mmHg ± 17.53 vs. 39.59mmHg ± 15.61; p=0.0540)

Conclusion
Our results indicated that methamphetamine users had echocardiogram findings with decreased ejection fraction and increased left ventricular mass index, end-diastolic volume index, and right ventricular systolic pressure consistent with worse dilated cardiomyopathy comparison to non-users.

Introduction
Methamphetamine activates the sympathetic system, resulting in tachycardia, hypertension, vasospasm/vasoconstriction, and myocardial wall stress or ischemia [1]. The exact prevalence of methamphetamine associated cardiomyopathy is unknown; however, the prevalence is on an incline due to increased drug usage. An echocardiogram is a noninvasive test commonly used to assess myocardial and valvular structure, quantify chamber size, and estimate ejection fraction. Additional specific details such as wall motion, wall thickness, and chamber size have been correlated to clinical conditions. Methamphetamine associated cardiomyopathy has been reported to have typical echocardiographic findings of dilated cardiomyopathy with reduced left ventricular systolic function and cardiac chamber enlargement [1]. Many overlapping features may be seen on echocardiograms from the various etiologies of cardiomyopathy, such as ischemic heart disease [2]. Assessments for wall motion abnormalities, cardiac chamber enlargement, and decline in left ventricular ejection fraction provide information and prognosis about an individual patient’s clinical condition. Typical factors attributed to influence left ventricle mass index (LVMI) include body size, ethnicity, and exercise-related factors. However, LVMI has been shown to

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predict cardiovascular events and premature death independently \cite{3,4}. Other parameters such as left ventricular end-diastolic volume (LV EDV) indicate left ventricular function and are standardized to a body-surface-area ratio. It represents the volume of blood at the end load filling and can be quantified via echocardiogram. Depending on the value, it may indicate enlargement compared to a normal value of <82 ml/m$^2$ \cite{5}. The right ventricle is designed to deliver a venous return to a low-pressure system. Right ventricular systolic pressure (RVSP) has been adopted as a marker to evaluate for pulmonary hypertension. It is equivalent to pulmonary artery systolic pressure in the absence of pulmonary outflow tract obstruction and is associated with reduced survival in patients with heart disease \cite{6}. The purpose of our study was to characterize various echocardiographic findings, including ejection fraction, right ventricular systolic pressure, cardiac mass index, and left ventricular end-diastolic volume amongst heart failure patients with and without a history of methamphetamine use.

**Materials And Methods**

We intended to perform a case-control study in heart failure patients (HF) patients with active methamphetamine usage compared to patients with cardiomyopathy without methamphetamine use. We observed the echocardiographic characteristics in both patient populations. The primary aim of this study is to determine whether there is any significant difference in echocardiographic parameters in heart failure patients with a history of methamphetamine use that would potentially determine their prognosis compared to patients without methamphetamine use. We included Right Ventricular Systolic Pressure (RVSP), Left Ventricular Ejection Fraction (LVEF), Left Ventricular Mass Index (LVMI), End Diastolic Volume Index (EDVI), grade of Diastolic Dysfunction (DD) along with age and gender to determine if there is any statistical difference between the two groups. The ejection fraction was calculated using the Modified Simpson Method (biplane method of disks).

After obtaining institutional review board (IRB) approval, we performed a retrospective chart review. We screened patient charts from November 1, 2019, to November 1, 2020, using heart failure related International Classification of Diseases, Tenth Revision, Clinical Modification (ICD-10 CM) codes that showed 1410 records. Our inclusion criteria included patients aged 18 years and above with clinical heart failure and active and recent methamphetamine use (in the last six months) based on history provided or urine toxicology, along with patients who underwent echocardiography during the index admission. We excluded patients who either left the hospital or expired before an echocardiogram was obtained. The final case group included 254 patients (Figure 1). Controls were screened using the same criteria except that patients were negative for methamphetamine on urine toxicology and history. Charts were also reviewed for a history of methamphetamine use. Patients were included in the case group if they were actively using methamphetamine based on history, even if urine toxicology was negative.
FIGURE 1: Selection criteria for the study

Data was collected for index admission, defined as the first admission to our institution for heart failure symptoms. The parameters were taken from echocardiography that was performed in our institution. The results for RVSP, LVMI, EDVI are reported as a continuous numerical value. As for LVEF, it is either reported as an exact numerical value or as a range within five value points. As a result, LVEF was rounded off to the nearest numerical value or an average value was taken for the range as follows: LVEF of <10 was entered as 10; 10-15 as 12; 15-20 as 17; 20-25 as 23; 25-30 as 27; 30-35 as 33; 35-40 as 37; 40-45 as 43; 45-50 as 47; 50-55 as 52; 55-60 as 58. Diastolic dysfunction is reported as grade 1 to 3. Grade 0 was used for patients who had no diastolic dysfunction.

All the data was compiled into an excel sheet. Patient identifiers were not disclosed except to the members of the data collection team. Data were analyzed using R version 4.0.0. A two-sample one-tailed t-test was used for RVSP, LVEF, LVMI, EDVI and age. Odds ratio and chi-square test of independence were used for diastolic dysfunction. Subgroup analyses were performed for parameters where the values were not available for all the patients.

Results

The final case group included 254 patients after the application of inclusion and exclusion criteria. The final control group included 268 patients. Majority were males, 178 (70%) and 180 (67%) in the case and control group respectively which was not statistically significant (odds ratio = 1.1450, p = 0.4735) (Table 1). The results for RVSP, LVEF, LVMI, EDVI and age are based on a two-sample one-tailed t-test, whereas diastolic dysfunction was reported using odds ratio and chi-squared test of independence.
Age was found to be statistically significant with the younger population in the case group (p = 0.0000). The mean value LVEF in the control group was higher than in the case group (p= 0.0000), suggesting better left ventricular systolic function in patients without methamphetamine use. LVMI, which signifies the mass of left ventricle as per body surface area, was significantly higher in the case group (p= 0.0000). So was the EDV Index (p=0.0000), signifying greater left ventricular dilatation in methamphetamine users. In patients where RVSP value was available (179 patients, 211 controls), mild statistical significance (p-value = 0.0540) was observed in the difference in RVSP between case and control groups (cases having higher RVSP). Given that not all the patients had RVSP reported, results may have been impacted by a smaller sample size (Table 2, Figures 2-6).

**TABLE 1: Case/Control by Gender**

|           | Male (n) | Female (n) | Odds ratio (95% CI); p-value |
|-----------|----------|------------|-----------------------------|
| Case      | 178      | 76         | 1.1450 (0.7907,1.6582); 0.4735 |
| Control   | 180      | 88         |                             |

**TABLE 2: Case/Control by Age**

| Age (Years) | n | Mean ± SD   | Median | SE | p-value |
|-------------|---|-------------|--------|----|---------|
| Case        | 254 | 50.74 ± 10.37 | 52     | 0.6507 | 0.0000 |
| Control     | 268 | 59.88 ± 13.97 | 61     | 0.8534 |

| LVEF (%)    | n | Mean ± SD   | Median | SE | p-value |
|-------------|---|-------------|--------|----|---------|
| Case        | 251 | 33.65 ± 18.02 | 33     | 1.1374 |
| Control     | 264 | 41.55 ± 15.61 | 43     | 0.9607 |

| LV Mass Index (grams/m²) | n | Mean ± SD   | Median | SE | p-value |
|--------------------------|---|-------------|--------|----|---------|
| Case                     | 190 | 122.49 ± 40.66 | 121    | 2.9498 |
| Control                  | 213 | 108.62 ± 32.82 | 105    | 2.2488 |

| EDV Index (mL/m²)        | n | Mean ± SD   | Median | SE | p-value |
|--------------------------|---|-------------|--------|----|---------|
| Case                     | 243 | 85.91 ± 37.40 | 83     | 2.3992 |
| Control                  | 257 | 72.44 ± 25.44 | 69     | 1.5869 |

| RVSP (mmHg)              | n | Mean ± SD   | Median | SE | p-value |
|--------------------------|---|-------------|--------|----|---------|
| Case                     | 179 | 42.29 ± 17.53 | 39     | 1.3103 |
| Control                  | 211 | 39.59 ± 15.61 | 36     | 1.0746 |

Abbreviations: Right Ventricular Systolic Pressure (RVSP), Left Ventricular Ejection Fraction (LVEF), Left Ventricular Mass Index (LVMI), End Diastolic Volume Index (EDVI), SD: Standard Deviation, SE: Standard Error, n: sample size
FIGURE 2: Box and Whisker plot for Case/Control by Age
FIGURE 3: Box and Whisker plot for Ejection Fraction (EF) in Case/Control Groups
FIGURE 4: Box and Whisker plot for Left Ventricular Mass Index (LVMI) for Case/Control groups
FIGURE 5: Box and Whisker plot for End Diastolic Volume Index (EDVI) for Case/Control groups
While comparing diastolic dysfunction between cases (32/129, 24.81%) and controls (35/134, 26.12%) where
dysfunction was defined as any grade (1, 2 or 3); the odds ratio was 1.07, not statistically significant (p =
0.807) which means that the incidence of diastolic dysfunction was similar in both groups. Diastolic
dysfunction was also broken down as per the grade, which again showed no statistical significance (p = 0.77),
demonstrating similar incidence between two groups regarding different dysfunction grades. However, the
sample size was smaller in individual grade groups (Table 3).
TABLE 3: Case/Control by Diastolic Dysfunction

| Diastolic dysfunction   | Cases (n) | Controls (n) | Odds ratio (95% CI); p-value |
|-------------------------|-----------|--------------|-------------------------------|
| None (0)                | 32        | 35           | 1.07 (0.6151,1.8671); 0.807   |
| Any level (1/2/3)       | 97        | 99           |                               |
| Grade 1                 | 30        | 30           |                               |
| Grade 2                 | 41        | 48           |                               |
| Grade 3                 | 26        | 21           | p-value                       |

Discussion

Methamphetamine is a highly potent and addictive substance that was first created in the late 1920s to mimic the nasal vasoconstrictor ephedrine. The addition of the “methyl” group to amphetamines further enhanced its potency by making it lipophilic and its ability to penetrate the blood-brain barrier. In chronic methamphetamine users, the drug permanently alters the user’s neurological function by affecting cognitive, psychiatric, and behavioural modalities. Similarly, methamphetamine users have various cardiovascular manifestations that include but are not limited to accelerated coronary plaque formation, cardiac arrhythmias, pulmonary hypertension, coronary vasospasm, and cardiac remodelling leading to cardiomyopathy, hypertension, aortic dissection, and acute coronary syndromes (Figure 7). Due to this, cardiovascular disease is the second leading cause of death in methamphetamine users after overdose [7].
Although methamphetamine consumption occurs independent of socioeconomic status, geographics, race, and culture, most consumers tend to be younger. The cardiac complications of methamphetamine are hypothesized to arise from a variety of mechanisms. In human autopsy specimens, severe interstitial fibrosis and scar formation has been documented [8]. Physiological pathways implicated include catecholamine surges acutely during methamphetamine intoxication with contingent hypertensive crises, longer-term upregulation of the sympathetic axis, and myocardial toxicity with impaired cellular metabolism. Depending on which process predominates, different patterns of pathology may develop, particularly in the case of methamphetamine-associated cardiomyopathy [9].

Preclinical studies have also shown that methamphetamine-induced endothelial nitric oxide synthase activation and endothelin-1 release leads to potent vasoconstriction [10]. Further, chronic methamphetamine users are less responsive to the vasodilatation effects of nitroglycerin [11].
Prolonged use of methamphetamine, especially when used intravenously, directly accumulates in the lungs. It is then hypothesized to be engulfed by pneumocytes leading to a similar surge of radical oxygen species formation, causing endothelial damage and pulmonary hypertension. Methamphetamine users are at a 27% increased risk of sudden cardiac death, as reported by the United States National Inpatient Sample database that studied over 180,000 methamphetamine drug users. Autopsies obtained from chronic methamphetamine users showed cardiac fibrosis and necrosis findings that were directly proportional to the duration and frequency of drug use. These chronic structural changes likely increased the risk of ventricular arrhythmia due to electrical conduction abnormalities of the heart [1,7].

Methamphetamine causes systolic dysfunction by causing chronic remodelling and left ventricular dilatation. In a study performed by Yu et al., mice were administered large doses of methamphetamine and 12 weeks later were found to have concentric left ventricular hypertrophy, myocardial fibrosis, necrosis and decreased contractile capacity [12]. In contrast, in an additional study performed by Lord et al., echocardiographic parameters were studied in mice given binge doses of methamphetamine. This study showed left ventricular dilatation with contractility and relaxation dysfunction - and, subsequently, an increase in left ventricular volume and a decrease in left ventricular wall thickness leading to eccentric dilated cardiomyopathy. Impaired relaxation suggested diastolic dysfunction with echocardiogram findings of increased end-diastolic volume and pressure [13].

A similar study performed in Germany by Schurer et al. studied thirty endomyocardial biopsies of methamphetamine users to evaluate their overall outcome with or without cessation of methamphetamine, the extent of fibrosis affecting clinical symptoms, and echocardiographic features. Their results were significant for the echocardiographic prevalence of impaired ejection fraction, increase in left ventricular end-diastolic diameter, with improvement in these factors with discontinuation of the drug [14].

To the best of our knowledge, there is only one previous study before ours that studied these specific echocardiogram features in methamphetamine users: ejection fraction (EF), end-diastolic volume (EDV), left ventricle mass index (LVMI), right ventricle systolic pressure (RVSP). In this study performed by Ito et al., in Hawaii in 2009, 28 methamphetamine users (case group) were compared to non-users revealing a statistically significant decrease in LVEF, increase in LV EDV; however, the results for LVMI and RVSP were not statistically significant. This possibly was due to their small sample size. Similar to cocaine, methamphetamine is hypothesized to activate the calcium/calmodulin-dependent protein kinase pathway that causes myocardial fibrosis and necrosis; methamphetamine likely cause the same effect physiologically. Though this study did not show a statistically significant increase in LVMI, they acknowledged their limitations due to the small sample size. Their results were still consistent with autopsy studies performed in San Francisco, California, that supported evidence of methamphetamine users to have enlarged hearts that weighed more than their age-matched controls. Our study showed statistically significant results for an increase in LVMI in methamphetamine users, which we hypothesize due to a chronic burden of hypertension and tachycardia leading to LV hypertrophy. The exact mechanism of myocardial injury remains unknown; however, a combination of LV hypertrophy due to significant vasoconstriction increased myocardial afterload and myocardial necrosis, fibrosis leading to eccentric dilatation and impaired relaxation causes this distinctive catastrophic heart failure [15].

Cocaine abuse causes higher LVMI, and lower EF [16,17]. Methamphetamine like cocaine has sympathomimetic effects causing vasoconstriction of coronary arteries, tachycardia, and direct myocardial toxicity; both may also have similar pathophysiological effects on the myocardium. Indeed, the sympathomimetic and toxic effects on the heart induced by methamphetamine are likely worse than cocaine because it increases catecholamine release rather than just inhibiting its reuptake [18].

Our study is limited by several factors but mainly by the inclusion of only select echocardiographic parameters. We also did not include any demographic data other than age and gender. The small sample size also limits our study as we only looked into patients’ data spanning one year (from 2019 to 2020) due to a change in our electronic medical record database before the year 2019. Our case group included patients with heart failure that had recent or active methamphetamine usage as correlating factor, not necessarily as the only causation of heart failure.

Conclusions
In conclusion, patients with methamphetamine-related cardiomyopathy tend to be younger and had worse cardiomyopathy related echocardiogram parameters when compared to patients with cardiomyopathy without methamphetamine use. Methamphetamine users had increased LV mass index, worse systolic dysfunction and increased chances of developing right heart failure from increased overall RVSP. Further large scale, randomized controlled trials are needed to confirm these findings of methamphetamine-related cardiomyopathy.

Additional Information
Disclosures
Human subjects: Consent was obtained or waived by all participants in this study. Kern Medical IRB issued approval 20090. Date Issued: December 22, 2020 Janpreet Singh Bhandohal, MD, Principal Investigator Department of Medicine, Division of Infectious Disease Kern Medical 1700 Mt. Vernon Avenue Bakersfield, CA 93306 Study # 20090: "Methamphetamine related cardiomyopathy: Characteristics and Outcomes" Dear Dr. Bhandohal, ACTION: The proposed project, named above, is appropriate for Expedited Review (45 Code of Federal Regulations 46, "Research Which Has Been Approved for Expedited Review", Category 5). The standard applies to requests for use of patient records that have been, or will be, created in the course of providing service to patients. Review using this standard requires that all database content be de-identified; none of the HIPAA PHI identifiers may be used for data. HIPAA research review includes authorization or waiver for data access, privacy for identifier use, data use limited to essential data for the case report description and identity security. HIPAA privacy regulations also require that publication/presentation subject information be de-identified. ☐ For the study listed above, the study plan, bibliography, data request, waiver of consent with protection for HIPAA-specified individual identifiers/study data, and templates for the database and identity protection are included in this review. ☐ To meet the required identity protections, the Principal Investigator is responsible to provide for subject identity protection: identity electronic storage retention on a Kern Medical-supported computer in a password protected file with access limited to study team members ☐ Subject identity cannot leave Kern Medical by any means, including electronic, digital, paper, video or voice. ☐ The study team members eligible to participate in the above-named study include: Principal Investigator: Janpreet Singh Bhandohal, MD Sub-investigators Vishal Kumar Narang, MD Roopam Jariwal, MD Michael Valdez, MD Nadia Raza, MD Leila Moosavi, MD Ramanjeet Singh Sidhu, MD Shikha Mishra, MD Page 2 of 3 Baldeep Kaur Mann, MD Theingi Tiffany Win, MD Fowrooz Joolhar, MD Asian Ghandforoush, DO REVIEW OUTCOME & INVESTIGATOR RESPONSIBILITY FOR STUDY CONTINUATION Clinical records for patient care/data abstraction created between 01/01/2010 through 11/30/2020 received approval for research use. ☐ The waiver of consent for use of medical records that have been or will be created is approved. ☐ Should you wish to continue data collection beyond the end of the approval cycle, continuation review and IRB approval is necessary; please either apply for continuing review or file a closing report at least two weeks prior to the end date of the approval cycle. ☐ Please refer to "Research Records Storage, Retention and Destruction", COM-1M-550, for information about research records responsibilities. ☐ The study approval cycle is 12/22/2020 through 12/21/2021. The end date of the approval cycle is the end date for records access for data collection. CHANGES TO STUDY TEAM, PROTOCOL OR STUDY DOCUMENTS ☐ Should you, Dr. Bhandohal, wish to delete or add research-eligible members to your study team, notification to the IRB Office prior to revising study team member deletion/addition(s) is necessary. You are welcome to send your request, which you may conveniently communicate by email, to IRB staff. ☐ Study design, study document changes and study continuation after the last date of the approval cycle must have IRB review & approval of proposed changes/continuation prior to change initiation/continuation. IRB COMPLIANCE WITH FEDERAL REGULATIONS ☐ Validation of expedited approval by the full IRB will occur at a convened meeting of the IRB. The following is a listing of current members of the Institutional Review Board: Randolph Fok, MD, PhD, Chair Arash Heidari, MD, Vice Chair Kate Botner, MSL, CPC, CMT Shahzad Chaudhry, MS, LMFT Shannon Hochstein, JD Jeffrey Jolliff, PharmD, BCPS, BCACP, AAHIVP, CDE Marie Joy Quiton-Buaya, PsyD, LMFT Dana Brucker, RN, Alternate Scientific Member ATTACHMENTS: Waiver of Consent Approval, approved templates. Animal subjects: All authors have confirmed that this study did not involve animal subjects or tissue. Conflicts of interest: In compliance with the ICMJE uniform disclosure form, all authors declare the following: Payment/services info: All authors have declared that no financial support was received from any organization for the submitted work. Financial relationships: All authors have declared that they have no financial relationships at present or within the previous three years with any organizations that might have an interest in the submitted work. Other relationships: All authors have declared that there are no other relationships or activities that could appear to have influenced the submitted work.

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