**Background:** Hypereosinophilic syndrome is a hematologic disorder characterized by an elevated eosinophil count in the peripheral blood and eosinophilic infiltration of other organs, resulting in end organ damage \(^1,2\). Such eosinophilic infiltration of the heart may result in the development of a restrictive cardiomyopathy, leading to symptomatic heart failure.

**Case report:** The patient was 23 years old at the time of diagnosis, previously a healthy young man. His initial complaint was exertional dyspnea, paroxysmal nocturnal dyspnea and peripheral edema. Laboratory findings revealed a significantly elevated eosinophil count (41% of the total number of leukocytes), as well as elevated levels of NT-proBNP (3784 pg/mL). He was then referred to our center from a local hospital. Initial echocardiographic examination revealed a slightly dilated left ventricle, with signs of a restrictive cardiomyopathy. Apical 4 chamber view revealed that the apex of the left ventricle is obliterated by a substance, suspected a thrombus formation (Figure 1). Magnetic resonance performed with gadolinium contrast revealed diffuse fibrosis of the endocardium and a severely hypokinetic apex of the left ventricle, filled with a thrombus mass (dimensions: 2.7x2.5cm), concluding that the finding is characteristic for endocardial fibroelastosis. In coordination with a hematologist, the patient was started on imatinib, for the eosinophilia, and low molecular weight heparin, as well as optimal medical treatment for heart failure. During one of the scheduled checkups, a follow up echocardiographic examination revealed a significant reduction in the overall size of the thrombus mass, but also a peduncular formation protruding into the ventricle (Figure 2). At that time, the patient had exhibited a significant clinical improvement, and was regularly

**KEYWORDS:** hypereosinophilic syndrome, cardiac thrombus, echocardiography.

**CITATION:** Cardiol Croat. 2017;12(4):139–140. | https://doi.org/10.15836/ccar2017.139

**ADDRESS FOR CORRESPONDENCE:** Nina Jakuš, Klinički bolnički centar Zagreb, Kišpatićeva 12, HR-10000 Zagreb, Croatia. / Phone: +385-91-5605-795 / E-mail: nina.jakush@gmail.com

**ORCID:** Nina Jakuš, http://orcid.org/0000-0001-7304-1127 • Ivo Planinc, http://orcid.org/0000-0003-0561-6704 • Hrvoje Jurin, http://orcid.org/0000-0002-2599-553X • Marijan Pašalić, http://orcid.org/0000-0002-3197-2190 • Dora Fabijanović, http://orcid.org/0000-0003-2633-3439 • Daniel Lovrić, http://orcid.org/0000-0002-5052-6559 • Jure Samardžić, http://orcid.org/0000-0001-7177-2206 • Boško Skorić, http://orcid.org/0000-0001-5979-2346 • Jana Ljubas Maček, http://orcid.org/0000-0001-7285-4026 • Bojan Biočina, http://orcid.org/0000-0001-9101-1570

**FIGURE 1.** Initial echocardiogram indicating the existence of a thrombus formation in the apex of the left ventricle.

**FIGURE 2.** Follow up echocardiogram performed after 7 months of treatment, showing a significant reduction in the overall size of the thrombus mass, but also a peduncular formation protruding into the ventricle.
Therapeutic challenges in treatment of a restrictive cardiomyopathy due to hypereosinophilic syndrome in a young patient: a case report

undertaking physical activity. Due to a significant embolization risk, he was urgently referred to cardiac surgery, where a successful surgical excision of the thrombus formation was performed, through a medial sternotomy.

**Conclusion:** Thrombus formation in patients with restrictive cardiomyopathy due to hypereosinophilic syndrome has been previously described in literature. On the other hand, complications of treating such thrombus mass are rarely described. This case report stresses the need for close echocardiographic monitoring during the dissolution of such large thrombus masses, anticipating potential embolic complications.

**LITERATURE**

