

Supplementary material

Redzak A, Preveden A, Todic M. Primary neuroendocrine tumor of the heart. Successful management of an extremely rare disease. Kardiol Pol. 2022.

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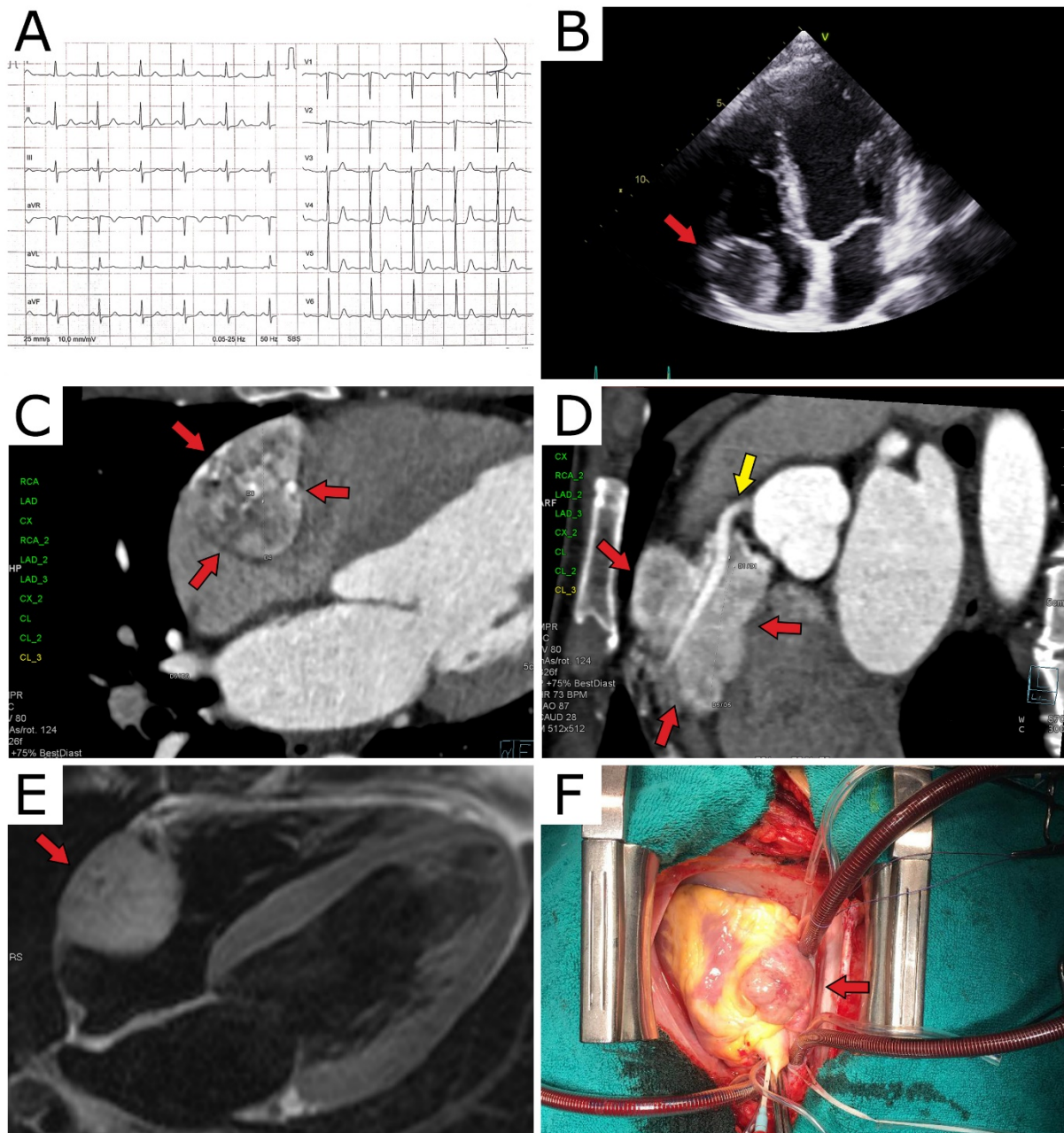


Figure S1. A. Tumor gross appearance after the excision: the tissue is brown and focally permeated with blood on serial sections. B. Pathohistology finding: uniform polygonal tumor cells, round nuclei, fine-grained and diffusely distributed chromatin, pale eosinophilic

cytoplasm arranged in organoid, trabecular and pseudoglandular formations, HE 40×. **C–F.** Immunohistochemical analyses: positive tumor cells in Ki67, CD56, synaptophysin and chromogranin, 20×

Video S1. Transthoracic echocardiography, apical 4-chamber view: tumor is located near the right atrium and collapsing it