

# Correction to: ASNC/AHA/ASE/EANM/HFSA/ISA/SCMR/SNMMI expert consensus recommendations for multimodality imaging in cardiac amyloidosis: Part 1 of 2—evidence base and standardized methods of imaging

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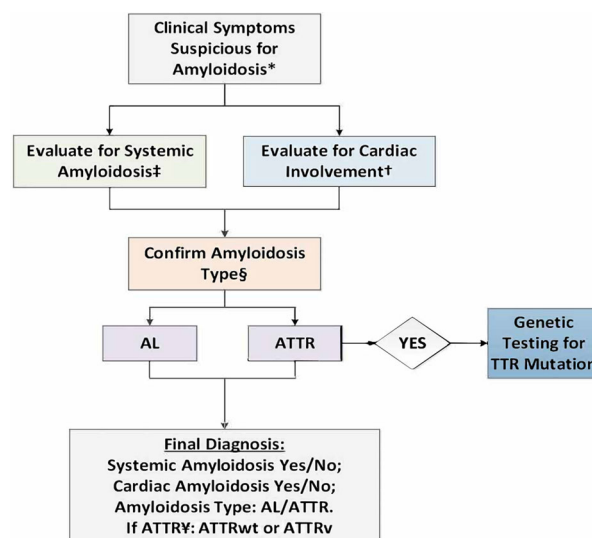
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## CORRECTION TO:

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- In the **Introduction** SCMR was listed incorrectly. SCMR is the Society for Cardiovascular Magnetic Resonance.
- **Figure 1** erroneously printed without Yen sign (¥) in 'Final Diagnosis.' Please see revised Figure 1.
- **Acknowledgments** erroneously printed without reviewers Richard Cheng, MD and Roy John, MD.

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**Figure 1. Systematic evaluation of cardiac amyloidosis.** A comprehensive evaluation of cardiac amyloidosis includes consideration of clinical symptoms, evaluation of cardiac involvement (biomarkers and cardiac imaging), evaluation of systemic amyloidosis (serum, urine testing, and biopsy), followed by typing of amyloid deposits into AL or ATTR, and documentation of mutations in patients with ATTR amyloidosis. **\*Clinical symptoms:** heart failure, peripheral/autonomic neuropathy, macroglossia, carpal tunnel syndrome, periorbital bruising, stroke, atrial fibrillation, postural hypotension, fatigue, weight loss, pedal edema, renal dysfunction, diarrhea, constipation. **†Evaluation for cardiac amyloidosis:** ECG, ECHO, CMR, EMB, <sup>99m</sup>Tc-PYP/DPD/HMDP/<sup>123</sup>I-mIBG/PET, NT-proBNP, troponin T. **‡Evaluation for systemic amyloidosis:** AL: detect plasma cell clone: serum and urine immunofixation, serum FLC assay and immunoglobulin analysis; AL: detect systemic organ involvement: 24-hour urine protein, alkaline phosphatase, eGFR, cardiac biomarkers (NT-proBNP, troponins); Tissue biopsy: EMB/Fatpad/Bone marrow/Other with Congo red staining. **§Confirm Amyloidosis Type:** ATTR: IHC and MS of Biopsy or <sup>99m</sup>Tc-PYP/DPD/HMDP Grade 2 or 3 if a clonal process is excluded; AL: MS or IHC of Biopsy. **¥Confirm TTR mutation in patients with ATTR amyloidosis:** genetic testing for TTR mutations. **AL**, amyloid light chain; **ATTR**, amyloid transthyretin; **CMR**, cardiac magnetic resonance imaging; **DPD**, -3,3-diphosphono-1,2-propanodicarboxylic acid; **ECG**, electrocardiogram; **EMB**, endomyocardial biopsy; **ECHO**, echocardiogram; **eGFR**, estimated glomerular filtration rate; **HMDP**, hydroxymethylenediphosphonate; **IHC**, immunohistochemistry; **mIBG**, meta-iodobenzylguanidine; **MS**, mass spectroscopy; **v**, hereditary; **PYP**, pyrophosphate; **Tc**, technetium; **wt**, wild-type.