

# A simplified diagnostic algorithm for adults with suspected systemic mastocytosis (SM)

This simplified diagnostic algorithm is based on the ECNM User's Guide (Valent 2022 *et al.*)<sup>1</sup> and other supporting published expert opinion (Theoharides *et al.*)<sup>2</sup> and is not intended to be a diagnostic tool. It does not replace the need for a complete evaluation of the patient by a healthcare professional.

**Common symptoms that may trigger suspicion of SM include characteristic skin lesions, unexplained osteoporosis, anaphylaxis, histamine-induced symptoms (cramping, headache, hypotension, diarrhoea), cytopenia, eosinophilia and splenomegaly<sup>1</sup>**

Serum tryptase testing should be performed upon clinical suspicion of SM<sup>1,2\*</sup>

**Serum tryptase <11.5 ng/mL<sup>2</sup>**

Mastocytosis unlikely – but cannot be ruled out<sup>1,2</sup>

**Follow up or monitor for increasing tryptase levels or clinical symptoms suggestive of SM<sup>1,4</sup>**

**Serum tryptase 11.5–20 ng/mL<sup>2</sup>**

SM possible – screen for *KIT* D816V peripheral blood with high-sensitivity assay<sup>1,2</sup>

**Serum tryptase ≥20 ng/mL<sup>2</sup>**

WHO minor criterion<sup>3</sup>

Conduct a bone marrow biopsy or screen for *KIT* D816V peripheral blood with high-sensitivity assay<sup>1,2</sup>

**Screen for *KIT* D816V mutation with droplet digital polymerase chain reaction (ddPCR)<sup>1,2</sup>**

**A highly sensitive PCR assay (e.g. ddPCR with ~0.01% sensitivity) is recommended for screening *KIT* D816V in peripheral blood.** Next-generation sequencing (NGS) assays have low sensitivity (~1–5%) and are not sufficient to detect *KIT* D816V mutation<sup>5</sup>

In a study of ISM patients (n=39):<sup>6</sup>

**95%**

detection of *KIT* D816V mutation with ddPCR assay

**28%**

detection of *KIT* D816V mutation with NGS assay

**Further work-up involves assessing if the patient meets the WHO diagnostic criteria (see the inside pages for full WHO criteria)**

**How to investigate an SM diagnosis**

Use this guide to help confirm an SM suspicion


\*Serum tryptase should be measured when the patient is at baseline and not immediately after an anaphylactic or mast cell activation event.<sup>4</sup>

ECNM=European Competence Network on Mastocytosis; ISM=indolent systemic mastocytosis; WHO=World Health Organization.

# WHO criteria are used to diagnose SM<sup>3</sup>


Diagnosis of SM requires 1 major and ≥1 minor criterion OR ≥3 minor criteria

Major criterion




**Multifocal dense infiltrates of mast cells (≥15 mast cells in aggregates)**  
In bone marrow biopsies and/or in sections of other extracutaneous organ(s)


Minor criteria




**≥25% of all mast cells are atypical cells (type I or type II) on bone marrow smears or are spindle-shaped in mast cell infiltrates**  
Detected in sections of bone marrow or other extracutaneous organs



**Activating KIT point mutation(s) at codon 816 or in other critical regions of KIT**  
In bone marrow or another extracutaneous organ



**Mast cells in bone marrow, blood or another extracutaneous organ express one or more of: CD2 and/or CD25 and/or CD30**



**Baseline serum tryptase concentration >20ng/mL (in the absence of an unrelated myeloid neoplasm)**  
In the case of a known HaT, the tryptase level should be adjusted

# WHO B- and C-findings are used to determine the subtype of SM<sup>3</sup>

C-findings

- Cytopenia/s:** ANC <1×10<sup>9</sup>/L, Hb <10g/dL, PLT <100×10<sup>9</sup>/L (one or more found)
- Hepatopathy:** Ascites and elevated liver enzymes± hepatomegaly or cirrhotic liver ± portal hypertension
- Spleen:** Palpable splenomegaly with hypersplenism ± weight loss ± hypoalbuminemia
- Gastrointestinal tract:** Malabsorption with hypoalbuminemia ± weight loss
- Bone:** Large-sized osteolysis (≥2cm) with pathologic fracture ± bone pain

B-findings

- High mast cell burden:** Infiltration grade (mast cells) in bone marrow ≥30% in histology (IHC) and/or serum tryptase ≥200ng/mL and/or KIT D816V VAF ≥10% in bone marrow or peripheral blood leukocytes
- Signs of myeloproliferation and/or myelodysplasia:** Hypercellular bone marrow with loss of fat cells and prominent myelopoiesis ± left shift and eosinophilia ± leukocytosis and eosinophilia and/or discrete signs of myelodysplasia (<10% neutrophils, erythrocytes and megakaryocytes)
- Organomegaly:** Palpable hepatomegaly without ascites or other signs of organ damage or/and palpable splenomegaly without hypersplenism and without weight loss or/and lymphadenopathy palpable or visceral LN-enlargement found in ULS or CT (>2cm)

\*Signs of myeloproliferation and/or myelodysplasia must be discrete and stable (neither disappear nor progress) and must not reach diagnostic criteria of an MPN, MDS or MPN/MDS, in which case the diagnosis changes to SM-AHN. The presence of a myeloid AHN excludes B-findings and SSM by definition.  
\*\*Alkaline phosphatase levels are typically elevated in patients with Advanced SM and SM-induced liver damage. In some of these patients, only elevated liver enzymes but no (clinically relevant) ascites are found.  
AHN=associated hematologic neoplasm; ANC=absolute neutrophil count; CT=computed tomography; Hb=haemoglobin; IHC=immunohistochemistry; LN=lymph node; MDS=myelodysplastic syndrome; MPN=myeloproliferative neoplasm; PLT=platelet count; SSM=smouldering systemic mastocytosis; ULS=ultrasound; VAF=variant allele frequency; WHO=World Health Organization.

# Diagnosing SM: An overview<sup>3,7</sup>

This simplified diagnostic algorithm is based on the WHO criteria<sup>3,7</sup>

