

Medical Al Ensemble Clinical Decision Report

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Primary Diagnostic Consensus

Diagnosis	ICD-10	Agreement	Confidence	Status
Familial Mediterranean Fever (FMF) Evidence: Mediterranean descent, Recurrent fever episodes with abdominal, chest, and joint pain, Family history of similar symptoms, Elevated CRP, ESR, WBC during attacks	E85.0	0.0%	Very Low	PRIMARY

Alternative & Minority Diagnoses

Diagnosis	ICD-10	Support	Туре
Periodic Fever Syndrome (other types) Evidence: Recurrent fever episodes with similar symptoms	E85.8	11.1%	Alternative (10-29%)
Systemic Juvenile Idiopathic Arthritis (sJIA) Evidence: Recurrent fever and joint pain	M08.2	11.1%	Alternative (10-29%)
Adult-Onset Still's Disease (AOSD) Evidence: Recurrent fever and joint pain	M35.3	7.4%	Minority (<10%)
TRAPS Syndrome (Tumor Necrosis Factor Receptor-Associated Periodic Syndrome) Evidence: Recurrent fever and inflammatory symptoms	M15.4	3.7%	Minority (<10%)
Periodic Fever, Aphthous Stomatitis, Pharyngitis, and Adenitis (PFAPA) Evidence: Recurrent fever and inflammatory symptoms	M15.4	3.7%	Minority (<10%)
Behçet's Disease	M35.2	0.0%	Minority (<10%)
Hereditary Periodic Fever Syndromes	E85.8	0.0%	Minority (<10%)
Hyperimmunoglobulinemia D Syndrome (HIDS)	E85.8	0.0%	Minority (<10%)
Mevalonate Kinase Deficiency (MKD)	E85.8	0.0%	Minority (<10%)
Cryopyrin-Associated Periodic Syndromes (CAPS)	E85.8	0.0%	Minority (<10%)

Analysis Overview

Models Queried: 7

Successful Responses: 7

Consensus Level: High

Total Cost: <\$0.01

Free Model Disclaimer: This analysis was generated using free Al models

Free models may provide suboptimal results. For improved accuracy and reliability, consider using premium models with an

API key.

Critical Decision Points & Evidence Synthesis

Critical Decision Points

Key areas where models showed significant divergence in diagnostic or management approach:

Evidence Synthesis & Clinical Correlation

Symptom-Diagnosis Correlation Matrix

Symptom	Familial	Periodic	Systemic	Adult-On	TRAPS Sy	PFAPA	Behçet's	Heredita
Recurrent fever	Strong	Moderate	Moderate	Moderate	-	-	-	-
Abdominal pain	Strong	-	-	-	-	-	-	-
Chest pain	Strong	-	-	-	-	-	-	-
Joint pain	Strong	-	Moderate	Moderate	-	-	-	-
Elevated inflam	Strong	-	-	-	-	-	-	-

Legend: +++ Strong association, ++ Moderate, + Weak, - Not typical

Diagnostic Decision Tree

Step	Action	If Positive	If Negative
1	MEFV Genetic Test	→ Confirm FMF, Start Colchicine	→ Proceed to Step 2
2	Extended Genetic Panel	ightarrow Alternative periodic fever	→ Proceed to Step 3
3	Autoimmune Workup	→ Consider SLE/Still's	→ Consider IBD
4	Inflammatory Markers	→ Monitor progression	→ Reassess diagnosis

Executive Summary

Case Description

A 28-year-old male of Mediterranean descent presents with:

- Recurrent episodes of fever lasting 1-3 days
- Severe abdominal pain during episodes
- Chest pain with breathing difficulties
- Joint pain affecting knees and ankles
- Family history: Father and paternal uncle have similar symptoms
- Episodes occur every 2-3 weeks
- Labs during attack: Elevated CRP, ESR, and WBC
- Between attacks: Completely asymptomatic

Patient reports episodes started in childhood around age 7. Recent genetic testing is pending.

Key Clinical Findings

- · Positive family history of similar episodes
- Recurrent fever episodes
- Severe abdominal pain with peritoneal signs
- Migratory arthritis affecting large joints
- Elevated inflammatory markers (CRP, ESR)

Primary Recommendations

- Consider Familial Mediterranean Fever (FMF) among differential diagnoses
- Obtain Genetic testing to evaluate for inherited conditions for diagnostic confirmation

Primary Diagnosis Clinical Summaries

■ Key Clinical Findings

Finding	Supporting Evidence	Clinical Reasoning	
Mediterranean descent	Clinical presentation	Key diagnostic indicator	
Recurrent fever episodes with abdominal, chest, and joint pain	Clinical presentation	Key diagnostic indicator	
Family history of similar symptoms	Clinical presentation	Key diagnostic indicator	
Elevated CRP, ESR, WBC during attacks	Clinical presentation	Key diagnostic indicator	
Asymptomatic between episodes	Clinical presentation	Key diagnostic indicator	

■ Recommended Tests

Test Name	Туре	Priority	Rationale
Genetic testing to evaluate for inherited conditions	Laboratory	Urgent	Diagnostic confirmation
Blood tests for inflammatory markers, antimicrobial sensitivities	Laboratory	Urgent	Diagnostic confirmation

■ Immediate Management

Intervention	Category	Urgency	Clinical Reasoning
Genetic testing to evaluate for inherited conditions	Medical	Immediate	Critical intervention
Blood tests for inflammatory markers, antimicrobial sensitivities	Medical	Immediate	Critical intervention

■ Medications

Medication	Dosage	Route/Frequency	Indication		
Colchicine	0.6 mg	oral / twice daily	Prophylaxis attacks	of	FMF

Diagnostic Landscape Analysis

Detailed Diagnostic Analysis

The ensemble analysis identified **Familial Mediterranean Fever (FMF)** as the primary diagnosis with limited consensus among 5 models.

Detailed Alternative Analysis

Diagnosis	Support	Key Evidence	Clinical Significance
Periodic Fever Syndrome (other types) Evidence: Recurrent fever episodes with similar symptoms	11.1%	3 models	Less likely
Systemic Juvenile Idiopathic Arthritis (sJIA) Evidence: Recurrent fever and joint pain	11.1%	3 models	Less likely
Adult-Onset Still's Disease (AOSD) Evidence: Recurrent fever and joint pain	7.4%	2 models	Unlikely
TRAPS Syndrome (Tumor Necrosis Factor Receptor-Associated Periodic Syndrome) Evidence: Recurrent fever and inflammatory symptoms	3.7%	1 models	Unlikely
Periodic Fever, Aphthous Stomatitis, Pharyngitis, and Adenitis (PFAPA) Evidence: Recurrent fever and inflammatory symptoms	3.7%	1 models	Unlikely
Behçet's Disease	0.0%	0 models	Unlikely
Hereditary Periodic Fever Syndromes	0.0%	0 models	Unlikely
Hyperimmunoglobulinemia D Syndrome (HIDS)	0.0%	0 models	Unlikely

Minority Opinions

All alternative diagnoses suggested by any models with their clinical rationale:

- Adult-Onset Still's Disease (AOSD) (ICD-10: Unknown) 7.4% agreement (2 models)
 - Supporting Models: Unknown, Unknown
- TRAPS Syndrome (Tumor Necrosis Factor Receptor-Associated Periodic Syndrome) (ICD-10: Unknown) 3.7% agreement (1 models)

Supporting Models: Unknown

• Periodic Fever, Aphthous Stomatitis, Pharyngitis, and Adenitis (PFAPA) (ICD-10: R50.9) - 3.7% agreement (1 models)

Supporting Models: Unknown

• Behçet's Disease (ICD-10: Unknown) - 0.0% agreement (0 models)

Supporting Models:

• Hereditary Periodic Fever Syndromes (ICD-10: R50.9) - 0.0% agreement (0 models)

Supporting Models:

• Hyperimmunoglobulinemia D Syndrome (HIDS) (ICD-10: Unknown) - 0.0% agreement (0 models)

Supporting Models:

• Mevalonate Kinase Deficiency (MKD) (ICD-10: Unknown) - 0.0% agreement (0 models)

Supporting Models:

• Cryopyrin-Associated Periodic Syndromes (CAPS) (ICD-10: Unknown) - 0.0% agreement (0 models)

Supporting Models:

Additional Diagnoses Considered:

• Periodic Fever Syndrome (other types) (ICD-10: E85.8) - 42.9% (3 models)

Evidence: Recurrent fever episodes with similar symptoms

• Systemic Juvenile Idiopathic Arthritis (sJIA) (ICD-10: M08.2) - 42.9% (3 models)

Evidence: Recurrent fever and joint pain

Management Strategies & Clinical Pathways

Immediate Actions Required

Priority	Action	Rationale	Consensus
1	Genetic testing to evaluate for inherited conditions	Clinical indication	50%
2	Blood tests for inflammatory markers, antimicrobial sensitivities	Clinical indication	50%

Recommended Diagnostic Tests

Test	Purpose	Priority	Timing
Genetic testing to evaluate for inherited conditions	Diagnostic confirmation	Routine	As indicated
Blood tests for inflammatory markers, antimicrobial sensitivities	Diagnostic confirmation	Routine	As indicated

Treatment Recommendations

Treatment recommendations pending diagnostic confirmation.

Model Diversity & Bias Analysis

Model Response Overview & Cost Analysis

Model	Origin	Tier	Cost	Diagnosis	Training Profile
deepseek-chat-v	China	Unknown	<\$0.01	Familial Mediterranean Fever	General
deepseek-r1	China	Unknown	<\$0.01	Familial Mediterranean Fever	General
gemma-3-12b-it	USA	Unknown	<\$0.01	Periodic Fever Syndrome (likely Familial Mediterranean Fever - FMF)	General
llama-3.2-3b-in	USA	Free	Free	Reactive Arthritis	General
mistral-7b-inst	France	Free	Free	Familial Mediterranean Fever	General
gpt-oss-20b	USA	Unknown	<\$0.01	Familial Mediterranean Fever	General
shisa-v2-llama3	Japan/USA	Free	Free	Familial Mediterranean Fever	General

^{**}Total Estimated Cost: <\$0.01**

Understanding Training Profiles

Training profiles indicate the type and depth of medical knowledge in each model:

Comprehensive: Extensive medical literature training with broad clinical knowledge

Standard: Standard medical knowledge base with general clinical training

Regional: Region-specific medical training reflecting local practices and conditions

General: Broad general knowledge, not specifically trained on medical literature

Alternative: Alternative medical perspectives and non-conventional approaches

Al Model Bias Analysis

Al model bias analysis is generated during orchestration (Step 2). This comprehensive analysis examines cultural, geographic, and training data biases across the Al models used.

Detailed Model Responses

Complete diagnostic assessments from each model:

1. deepseek-chat-v (China, Released: 2024-12-26)

Primary Diagnosis: Familial Mediterranean Fever (ICD-10: E85.0) - Confidence: 0.95 **Differential Diagnoses:**

- Periodic Fever Syndrome (other types) (ICD: E85.8) 0.7
- Systemic Juvenile Idiopathic Arthritis (ICD: M08.2) 0.4
- Inflammatory Bowel Disease (ICD: K50.9) 0.3

Key Clinical Findings:

- Mediterranean descent
- Recurrent self-limited febrile episodes
- Abdominal pain
- Chest pain with pleurisy

2. deepseek-r1 (China, Released: 2025-01-20)

3. gemma-3-12b-it (USA, Released: 2024-12-11)

Primary Diagnosis: Periodic Fever Syndrome (likely Familial Mediterranean Fever - FMF) (ICD-10:

M15.4) - Confidence: 0.85

Differential Diagnoses:

- TRAPS Syndrome (Tumor Necrosis Factor Receptor-Associated Periodic Syndrome) (ICD: M15.4) 0.3
- Adult-Onset Still's Disease (ICD: M35.3) 0.25
- Systemic Juvenile Idiopathic Arthritis (sJIA) (ICD: M08.0) 0.2

Key Clinical Findings:

- Recurrent fever episodes (1-3 days)
- Severe abdominal pain during episodes
- Chest pain with breathing difficulties
- Joint pain (knees and ankles)

4. Ilama-3.2-3b-in (USA, Released: 2024-09-25)

Primary Diagnosis: Reactive Arthritis (ICD-10: M35.4) - Confidence: 0.8

Differential Diagnoses:

- Ankylosing Spondylitis (ICD: M45.9) 0.6
- Gonococcal Arthritis (ICD: A04.0) 0.4
- Psoriatic Arthritis (ICD: M07.9) 0.5

Key Clinical Findings:

- Recurrent episodes of fever, abdominal pain, and joint pain
- Elevated CRP, ESR, and WBC during attacks
- · Family history of similar symptoms
- 5. mistral-7b-inst (France, Released: 2023-09-27)
- 6. gpt-oss-20b (USA, Released: 2024-05-13)
- 7. shisa-v2-llama3 (Japan/USA, Released: 2024-12-20)

Primary Diagnosis: Familial Mediterranean Fever (FMF) (ICD-10: E85.0) - Confidence: 0.9 **Differential Diagnoses:**

- Systemic Juvenile Idiopathic Arthritis (sJIA) (ICD: M08.2) 0.6
- Adult-Onset Still's Disease (AOSD) (ICD: M07.2) 0.5
- Periodic Fever, Aphthous Stomatitis, Pharyngitis, and Adenitis (PFAPA) Syndrome (ICD: K00.8) 0.4

Key Clinical Findings:

- Recurrent febrile episodes with peritonitis-like abdominal pain
- Pleuropericarditis (chest pain, breathing difficulties)
- Arthritis affecting knees and ankles
- Family history of similar symptoms (paternal lineage)