					Comparison Chart of Systemic Autoinflammatory Diseases (SAID) Involving Periodic Fevers																
	Cryopyrin-Asso Familial Cold Autoinflammatory Syndrome	ciated Periodic Sy Muckle-Wells Syndrome	Neonatal-Onset Multisystem Autoinflammatory Disease—aka Chronic Infantile Neurological	Schnitzler Syndrome	Pyrin Familial Mediterranean Fever	Protein Folding Tumour Necrosis Factor (TNF)- Associated Periodic Sydrome -aka Familial Hiber-	Hyperimmuno- globulinemia D with Periodic Fever Syndrome –aka Mevalonate	Mevalonate Aciduria Syndrome	Infla  Deficiency of Interleukin-1ß (IL-1ß) Receptor Antagonist	mmatory Bone Dis	Chronic Recurrent Multifocal Osteomyelitis—aka Synovitis, Acne, Pustulosis, Hyperosto-	Familial Psoriasis (PSORS2)—aka	Pyogenic Sterile Arthritis, Pyoderma Gangrenosum, & Acne Syndrome	Granulomatous  Juvenile Systemic Granulomatosis  —aka Blau syndrome, Pediatric Granulomatous Arthritis (PGA), Farly Opent Sargaida	NLRP12-Associated Periodic Fever Syndrome—aka Familal Cold Autoinflammatory Syndrome	Neutrophilic Dermatosis with Lipodystrophy & Flevated Temper-	•	Periodic Fever S Periodic Fever, Aphthous Stomatitis, Pharyngitis, & Cervical Adenitis (PFAPA)	Systemic-Onset	Macrophage Ac Adult-Onset Stills Disease—aka Adult Still's, Wissler- Fanconi Syndrome	Lymphohistiocy- tosis—aka Familial Erythrophagocytic
ACRONYM	FCAS	MWS	Cutaneous Articular Syndrome (CINCA) NOMID/CINCA		FMF	nian Fever TRAPS	Kinase Deficiency HIDS/MKD	MA	DIRA	MAJEED		CAMPS/PSORS2		sis, or Jabs Syndrome	NLRP12/FCAS2	Nishimura Syndrome  CANDLE/JMP	BEHÇETS	-aka Marshall Syndrome PFAPA	soJIA/sJIA	AOSD	Lymphohistiocytosis  1° HLH/ FHL
GENE	NLRP3	NLRP3	NLRP3	Currently unknown.	MEFV	TNFRSF1A	MVK	MVK	IL1RN	LPIN2	Currently unknown.	CARD14	PSTPIP1	NOD2	NLRP12	PSMB8 and some other Proteasome genes.	ERAP1 (with HLA-B51); also variants near: CCR1, KLRC4, STAT4 42	Currently unknown.	Currently unknown.  HLA-DRB1 in some pts w/European ancestry <sup>57</sup>	Currently unknown.	PRF1, STX11, STXBP2, MUNC13-4, RAB27A X link: SH2D1A, BIRC4
INHERITANCE	Autosomal Dominant. Large familial groups, some spontaneous mutations 1	Autosomal Dominant. Spontaneous mutations, some familial groups. <sup>1</sup>	Autosomal Dominant. Spontaneous mutations, few familial cases. <sup>1</sup>	Unknown.	Autosomal Recessive. Some cases are genedosage-dependent autosomal dominant. <sup>10</sup>	Autosomal Dominant. Spontaneous mutations, some familial groups. <sup>1</sup>	Autosomal recessive. Some cases with only one mutation found. <sup>33</sup>	Autosomal recessive.	Autosomal recessive.	Autosomal recessive.	Currently unknown.	Autosomal Dominant. Spontaneous mutations, some familial groups. <sup>23</sup>	Autosomal Dominant. Spontaneous mutations, some familial groups. <sup>29,30</sup>	Autosomal Dominant.	Autosomal Dominant. Spontaneous mutations, some familial groups. <sup>29,30</sup>	Autosomal recessive.	Complex.	Currently unknown.	Complex.	Currently unknown.	Autosomal recessive, but if X-linked: inheritance is dominant.
ETHNICITY	Affects all races, but many are of European descent. <sup>1</sup>	Affects all races, but many are of European descent. <sup>1</sup>	Any-present in all races.1	Affects all races, but most cases are in Europe. 13 More men than women are affected.	Turk, Armenian, Arab, Sephardic Jew, Italian. Most common inherited periodic fever syndrome	Affects all races. 2nd most common inherited I SAID (after FMF.) <sup>1</sup>	Mostly of Dutch descent or Northern European. <sup>1</sup>	, Mostly of Dutch descent, or Northern European. <sup>1</sup>	Carriers in 0.2% population of Newfoundland & 1.3% in Puerto Rico. Also Dutch Brazilian & Lebanese pts. 16	documented cases are of Middle Eastern	Affects all races, but the majority of patients have European ancestry; more	Asian ancestry. Pts. in US	Currently, the only docu- mented cases are from Europe, New Zealand & the USA. <sup>30</sup>	Affects all races.	Unknown. Current cases from Guadeloupe, Italy, Armenia 38,39	Caucasian, Hispanic & Japanese. <sup>27</sup>	Rare in the USA. More common in the Middle East, Asia & Japan. (Silk Road Route.) <sup>42,43</sup>	Affects all races. <sup>40</sup>	Affects all races. soJIA accounts for 10% of all JIA. <sup>45</sup>	Rare. Affects all races. <sup>44</sup>	Affects all races. 80% of African Americans, & 20% of pts. w/European decent have <i>PRF1</i> mutations. 47,48
FREQUENCY IN THE WORLD	most cases are from	1:1 million, maybe more. Some large family groups. <sup>5</sup> Frequency of CAPS in France is 1:360,000. <sup>55</sup>	Estimated frequency 1:1 million, mostly due to spontaneous genetic mutations. <sup>5</sup>	Unknown. Over 150 known cases, mostly in	In specific ethnic groups, the carrier frequency of <i>MEFV</i> variants is up to 1:5 people.	Unknown. TRAPS affects 0.01:10,000 people in the - European Union. <sup>51</sup> >1000	Unknown, but very rare. >200-300 known patients worldwide, (>300, when suspected cases are also included.) <sup>12</sup>	Unknown, but very rare. <100 known patients worldwide. <sup>11</sup>	Unknown, but very rare. In some regions of Aricibo, Puerto Rico w/ more DIRA carriers, DIRA may occur 1:6300.16	Unknown, but very rare. Very few documented cases at this time. <sup>18,53</sup>	<u>'</u>	Unknown, but rare.	Unknown, but rare.	Unknown, but rare.	Unknown, but rare.	Unknown, but rare.	Prevalence is 80- 370:100,000 people in Turkey, 10:100,000 in Japan & 0.6:100,000 in Yorkshire, UK. <sup>3</sup>	Unknown. Most commo non-infectious recurren fever disorder. <sup>40</sup>	n Uncommon. 0.4—0.9 cas- t es per 100,000 people, per year. <sup>46</sup>	France: estimated that 0.16:100,000 people have AOSD. AOSD affects more women than men.44	1° HLH affects 1:50,000
TIMING OF SYMPTOMS OR ATTACKS	12-24 hours, or longer. Onset of fever & flares is often 1-3 hours after exposure to cold or cooling temperatures. <sup>1</sup>	fever & symptoms are often triggered by cold	Continuous w/increased symptoms & fever during flares.¹ Chronic inflammation noted between flares.	12-36 hours. Rash is present first. Intermit- tent fevers, that often occur separately from the rash. <sup>13</sup>	12-72 hours. <sup>1,9</sup> Recurrent fever & flares can occur weekly, or only a few times a year.	weeks. <sup>1,9</sup>	3-7 days. Recurrent bouts of fever & flares every 2-12 weeks. <sup>1,9</sup> Some flares occur after vac- cines. <sup>9</sup>		Continuous inflammation from birth/fetal development. Untreated DIRA can lead to death	Flares last for a few days, with 1-4 exacerba- tions a month of high fevers, severe pain, & joint swelling. 18,53	At least 6 months w/ chronic/relapsing symptoms. Often 7-25 yrs. of symptoms. Many bone lesions heal	Continuous chronic pustular or plaque psoriasis, triggered by inflammatory stimuli. Some cases w/psoriatic arthritic <sup>23,24</sup> .	Early-onset, destructive, recurrent inflammation of the joints, skin & muscle. Flares often occur after mild injury, or injections. <sup>29</sup>	daily fevers, rash &	1-3, to up to 7-15 days of fevers 39–40°C, rash and pain. Onset after exposure to cold or cooling temperatures. <sup>38,39</sup>	ease flares. Inflammatory	ent in almost all patients during flares for around 10 days w/inflamation in	toms lasting 3-6 days,	times/day for >2 weeks,	last for <4 hours, recurring more than once a week, w/a maculopapular rash &	Fevers often > 39°C 1-2 times/day for >2 weeks, most often occurring in the evening w/arthralgia, rash & other symp- toms. 47,48
AGE OF ONSET	ent with symptoms later	Infancy, but a few present with symptoms later in childhood or adolescence. <sup>1</sup>	Neonatal/early infancy. Rash, symptoms, & abnormal labs are often present at birth. <sup>1,6</sup>	Most cases start in middle age, over 35-50 yrs. Youngest pt. was 13 yrs old. Symptoms start w/the rash. 13		Most first attacks by 3 yrs, & almost all begin by 20 yrs. of age; a few start later in life. <sup>9</sup>	symptoms in infancy.9	Most present with symptoms at birth, or in early infancy. Most have facial features noted at birth. <sup>11</sup>	birth, or as a neonate: pustular rash, bone	Most present with symp toms in infancy to early childhood, between 3 weeks to 2 years of age. <sup>15</sup>	1 ' 1 1 1 1 1 1 1 1	from infancy—childhood to adulthood w/pustular	First symptoms of arthritis develop by 1-10	Rash often develops by 4 months of age, fevers and other symptoms present by 4 yrs. of age. <sup>3</sup>	Neonatal/early infancy. Rash, fevers, symptoms, may be present at birth. <sup>38,39</sup>	Onset at birth or in infancy. Progressive damage from chronic inflammation noted as the child grows. <sup>26,27</sup>	Most show symptoms in early adulthood (20's-30's) but the onset can be in childhood, or any age. 42,43	Early childhood, usually between 2-5 years of age. A few adult-onset cases. Many teens out- grow it. <sup>40</sup>	16-most often by 2 years of age, or between 0-5	First onset of symptoms occurs between 16-35	Onset <1yr: often by 6 months—early childhood. Some in utero or late childhood. A few adult- onset cases. 47,48,49
SYSTEMIC FIN							D.C.	l Diff											D 1 51 4		400/
SKIN/ CUTANEOUS	Cold induced urticaria- like rash with increased neutrophils at the eccrine coils. <sup>4</sup> Almost daily rash—increases w/flares. <sup>1</sup>	Urticaria-like rash with increased neutrophils at the eccrine coils. <sup>4</sup> Most w/daily rash that increases w/flares. <sup>1</sup>	Ever-present <sup>1</sup> Urticaria- like rash with increased neutrophils at the eccrine coils. Rash increases with flares. <sup>4</sup>	Maculopapular rash, & plaques (sometimes itchy) on the chest & limbs. Dermis has neutrophillic infiltrate. Dermographism. <sup>13</sup>	erysipeloid erythema on the ankle–foot–be- low knee region–lasts 2-3 days during flares of symptoms. <sup>1</sup>	Migrating rash w/deep pain under rash areas. Severe pain follows the rash path from the trunk out to the limbs. <sup>9</sup>	rash. Some w/petechiae or purpura present. A few w/apthous ulcers. <sup>1,9</sup>	Diffuse maculopapular or morbilliform rash. Some w/petechiae or purpura present. A few w/apthous ulcers. <sup>1,9,11</sup>	pustules at hair folicles. Oral ulcers, pathergy, hyperkeratosis, acanthosis; high neutrophil infiltrate of dermis. 16,26	Most patients have inflammatory dermatosis, Sweet's syndrome, pustular skin lesions, psoriasis. Intra-epiderma neutrophils. 18,53	Some patients have acne, &/or pustulosis on the palms &/or soles of their extremities (w/ SAPHO). 23% w/psoriasis. 19,22,54	Generalized pustular psoriasis (can be severe), &/or plaque psoriasis. Sometimes nails are affected w/psoriasis. <sup>23, 24</sup>	Pathergy. Pyoderma gangrenosum ulcerative lesions, &/or severe cystic acne. Affected tissues w/high neutrophil infiltration. <sup>29</sup>	First symptom: scaly plaques. The rash starts on face, then on torso. Biopsies w/ non-caseating granulomatous dermatitis. <sup>34</sup>	Present during flares: Urticarial or malar rash <sup>39</sup> noted in some patients; some with buccal aph- thosis. <sup>38,39</sup>	Annular cutaneous plaques w/residual purpura. Lipodsystrophy first on face & around joints. Lips swell w/flares. Purple-red eyelids. <sup>26,27</sup>	Pathergy. Pseudofol- liculitis, erythema nodosum-like &/or acneiform nodules. 98% w/mouth ulcers, & 65% have genital ulcers. 43	Some have a rash with flares. Aphthous stomatitis, & pharygitis with exudate, (but no infection) is a classic finding. 40,41		Evanescent, salmon- pink, mildly pruritic maculopapular rash on the proximal limbs and trunk. <sup>44</sup>	40% w/transient maculopapular, nodular or purpuric skin rashes during bouts of high fever. Jaundice. <sup>47,48</sup>
NEUROLOGIC	Some have headaches, fatigue w/fever after cold exposure. Unknown if there are notable CNS affects at this time.1	Some have headaches, fatigue w/fever & flares. Uncommon to have many other CNS symptoms. <sup>1</sup>	tive impairments. Papille	Intermittent fevers can rise > 40°C. Chills are uncommon. Fatigue & headaches are common w/fevers. Temperature changes, stress & exercise can trigger flares. 13	meningitis is rare and can occur during flares but is never chronic.¹ Other neurological involvement is very		Headaches & fevers w/flares of symptoms are common. <sup>1,9</sup> More severe neurological symptoms are rarely present in HIDS. <sup>9</sup>	Fevers w/flares. Microcephaly, dolichocephaly, mental retardation, developmental delays, cerebellar ataxia, cerebellar atrophy & epilepsy often develop over time. 11	High fevers are not common, or noted in the neonatal period. Neurological complications are not common. A few cases of cerebral vasculitis noted. 16,26	days w/flares & severe pain. Other neurological	of patients during flares of CRMO. Other neuro-logical symptoms are not noted. Some w/impaired		Fevers can accompany flares of joint inflam- mation and pain. Other neurological symptoms are not noted. <sup>31</sup>	Intermittent-persistent daily fevers. Some have cranial neuropathies. 80% have vision damage & joint deformities is untreated. Some cases have peripheral nerves affected. <sup>34</sup>	Fevers 39–40°C myalgia, headaches with flares. Other neurological symp- toms are not noted. <sup>31</sup>	Aseptic meningitis & systemic inflammation. Growth delays—low height & weight. Developmental delays. <sup>26,27</sup>	20-40% have Neuro-Behçets w/headaches, aseptic meningitis or meningoencephalitis, seizures, hemiplegia, or cranial nerve palsies. Cerebral venous thrombosis w/high ICP noted. <sup>43</sup>	High fevers for 3-6 days, with chills & malaise. Some patients have headaches with flares. Other neurological symptoms are not noted. 41	High fevers > 39°C 1-2 times/day for >2 weeks. Other neurological symptoms are rare. A few cases w/seizures, meningismus, irritabil- ity & decreased level of consciousness. <sup>46</sup>	ing fevers, fatigue and myalgia with flares. Other neurological symptoms are very rarely seen. <sup>44</sup>	High fevers. Increased CSF protein. High ICP. Multifocal inflammation of the gray & white matter, intracranial bleeding, generalized atrophy or brain edema,
AUDITORY	Some pts have mild hearing loss—not cur- rently known if it's from	Many have increased sensorineural hearing loss, starting in adoles-	Many have increased sensorineural hearing loss, from infancy/child-	Uncommon. <sup>13</sup>	Uncommon—not be- lieved to be caused by a FMF disorder. <sup>1</sup>	Uncommon—not believed to be caused by TRAPS. <sup>1</sup>		Uncommon—not be- lieved to be caused by MA. <sup>1,9,11</sup>	Not noted. 15,16	Not noted. <sup>18,53</sup>	Not noted. <sup>19,21,22,54</sup>	Not seen. <sup>23, 24</sup>	Not noted. <sup>29,30,31</sup>	Not noted. <sup>34</sup>	Many have increased sensorineural hearing loss. 38,39	Some have frequent otitis &/or recurrent sinusitis. <sup>27</sup>	Not noted. 42,43	Not noted. <sup>40,41</sup>	Not noted to be from soJIA. <sup>45,46</sup>	Not noted. <sup>44</sup>	Seizures or coma. <sup>47</sup> Not noted. <sup>47,48,49</sup>
OPHTHALMIC	CAPS inflammation.¹  Conjunctivitis (non-infectious) during flares.¹	Conjunctivitis (non-infectious) dur- ing flares, <sup>1</sup> or corneal haze. <sup>26</sup>	Papilledema, uveitis, iritis, conjunctivitis. Some w/retinal scarring, corneal haze or vision loss. 6,26	Not noted. <sup>13</sup>	Very rare to uncommon.	.1 Conjunctivitis, & periorbital edema during flares.1,9	Very rare to uncommon.	racts, blue sclerae & ta-	tivitis can be caused by	Not noted. <sup>18,53</sup>	Some cases of uveitis. <sup>19</sup>	Not seen. <sup>23, 24</sup>	Not noted. <sup>29,30,31</sup>	Uveitis (some w/blind- ness) 50% w/cataracts, 1:3 pts. get 2° glaucoma, inflamed conjunctiva, lacrimal glands, retina 8 optic nerves. <sup>34</sup>	Not noted. <sup>38,39</sup>	flammation on the eye.) Conjunctivitis. Kerati-	Frequent anterior &/or posterior uveitis. Cataract, retinal vasculitis <30% risk for blindness. Papilledema w/CNS involvement 43	Not noted. <sup>40,41</sup>	Uveitis can be a complication from soJIA.46	Not noted. <sup>44</sup>	Blindness due to CNS inflammation. <sup>48</sup>
CARDIO- PULMONARY	Not noted. <sup>1</sup>	Rare. <sup>1</sup>	Some have clubbing of fingers. Some cases of pericardial effusions, or pericarditis. <sup>1</sup>	Not noted. <sup>13</sup>	45% have pleuritis, pain ful respiration, w/flares. Some w/pericarditis.1	Common, including pleurisy. <sup>1</sup>	Rare. <sup>1</sup>	Rare. <sup>1,11</sup>	Some with resp. distress. 1 case: Pulmonary hemo- siderosis & progressive interstitial fibrosis. 15,16,17	Not noted. <sup>18,53</sup>	Not common—some patients also have ANCA+ Vasculitis that can affect the lungs. <sup>18,54</sup>	Not noted. <sup>23, 24</sup>	Not noted. <sup>29,30,31</sup>	Some have atrial hypertension &/or pericarditis Some cases with lung involvement. <sup>34,35</sup>	Not noted. <sup>38,39</sup>	Clubbing of the fingers &/or toes. At risk for cardiac arrythmias & dilated cardiomyopa- thy <sup>26,27</sup>	Myocarditis, endocarditis w/aortic or mitral insuffi- ciency, arterial aneurysm, pulmonary embolism. <sup>43</sup>	Flares of fevers, stomatitis & pharygitis are not associated with respiratory illness. 40,41	pericarditis) is often	<25% have pleuritis, pericarditis (a few w/ tamponade.) Some myo- carditis, pleural effusions, ARDS <sup>44</sup>	High risk for respiratory infections triggering fevers, systemic inflammation & MAS. Edema. <sup>49</sup>
ABDOMINAL	Uncommon. <sup>1</sup>	Some have abdominal pain w/flares or other gastrointestinal issues.1	Nausea, vomiting & abdominal pain with flares, or with high CNS pressure. <sup>6</sup>	GI symptoms are uncommon. Enlarged liver &/or spleen is common. <sup>13</sup>		Peritonitis, diarrhea, & constipation w/flares.1	Extreme pain, vomiting & diarrhea w/flares. <sup>1,9</sup>	Enlarged liver &/or spleen. Cholestatic liver disease. Pain, vomiting & diarrhea w/flares. <sup>1,9,11</sup>	Rarely have GI issues.  Mouth ulcers, stomatitis, & failure to thrive are	Enlarged liver & chole- static jaundice in the neonatal period, but it is	Some patients also have inflammatory bowel diseases. <sup>19</sup>	Not noted. <sup>23,24</sup>	Some patients also have irritable bowel syndrome. <sup>29</sup>	Enlarged liver &/or spleen. Some w/ GI pair higher risk for kidney &/ or liver issues. <sup>34,35,36</sup>		Loose bowels with flares. Enlarged liver & abdomen. Delayed or slow growth. <sup>26,27</sup>	Ulcers from mouth to anus. Nausea, abdomi- nal pain, anorexia, diar- rhea, (may be bloody). <sup>43</sup>	Abdominal pain, diar- rhea often present with flares. <sup>40,41</sup>	Peritonitis rarely occurs 50% have an enlarged spleen, some w/an enlarged liver. <sup>46</sup>	. 50-75% w/enlarged liver, abnormal LFTs. 43% w/ enlarged spleen. Renal disease is rare.44	Liver disease is com- mon. High risk of death from multi-organ failure in 2+ months if untreated. <sup>49</sup>
LYMPHATIC	Not noted. <sup>1</sup>	Rarely noted. <sup>1</sup>	Some pts. with enlarged liver and/or spleen, many have large lymph	<20% w/lymphoma, IgM myeloma, or Walden- ströms. >45% w/enlarged lymph nodes. <sup>13</sup>	common, some have	Enlarged spleen common, some have enlarged lymph nodes. <sup>1</sup>	Enlarged cervical lymph nodes are common in children.¹ Some have	·	Enlarged liver and/or spleen is common. Risk of organ failure if		Vasculitis that can affect		Not noted. <sup>29,30,31</sup>	Enlarged liver &/or spleen, enlarged lymph nodes. <sup>34,35,36</sup>	Some patients with adenopathy. <sup>39</sup>	Enlarged liver, with elevated liver enzymes; enlarged lymph	Some w/enlarged liver	Cervical adenopathy during flares. <sup>40,41</sup>	Many w/generalized lymphadenopathy. Some w/mesenteric adenitis. <sup>45,46</sup>	Lymphadenopathy is common. Many w/	Lymphoma. Hemophago- cytosis—spleen/lymph nodes. Large liver/
JOINTS/BONES MUSCLES & CARTILAGE	Arthralgias, stiffness & swelling with flares.	Arthralgias, recurrent arthritis, stiffness & swelling with flares. <sup>1</sup>	Joint pain, knee valgus or varus. Some w/frontal bossing, saddleback nose, contractures, clubbing.¹<50% of patients knees have bony overgrowth. Short stature, growth delays failure to thrive, arthritis osteopenia noted.¹.26	80% have muscle, bone &/or joint pain; arthritis. Bone pain is most common in the iliac and tibia. <40% have bone lesions. Some w/osteocondensation &	Mono/Polyarthritis, oligoarthritis & clubbing are common. Ankle arthralgias are common Severe arthritis of the hip or ankle is rare.1	the state of the s	enlarged spleens.  Arthralgias common, symmetric polyarthritis frequently noted.1	Congenital defects are often noted: micro-cephaly, dolichocephaly, wide irregular fontanels, low set and posteriorly rotated ears, downslanted palpebral fissures. Hypotonia, myopathy, & failure to thrive are common. <sup>11</sup>	Joint swelling, severe bone pain. Bone biopsy shows no infection. Common: Balloon-like widening of the anterior rib ends, periosteal elevation along multiple long bones, multifocal osteolytic lesions. Other bones affected. <sup>16</sup>	Periarticular tender soft tissue swelling. Bone biopsy shows no infection. Early-onset Chronic Recurrent Multifocal Osteomyelitis (CRMO), periarticular tender soft tissue swelling, short stature, delayed bone age, contractures. 18		· ·	Episodic inflammatory arthritis, often to one joint at a time that doesn't resolve on it's own. Intermittent sterile pauciarticular, peripheral erosive arthritis. Joint damage & destruction can often develop from the arthritis. 29,30,31,32,55	Symmetrical chronic polyarthritis or oligoar-thritis of the wrists, knees, ankles w/a boggy appearance is usually caused by an exuberant tenosynovitis. <sup>34,35,36</sup>	Myalgia, arthralgia, fatigue & malaise w/flares. Permanent bone or joint damage not noted.39	Joint Contractures, muscle atrophy, pan- niculitis induced lipodystrophy, myositis, fatigue and malaise. Inflammed nose & ear cartilage (chondritis). Growth delays—low height & weight. <sup>26,27</sup>	45% have arthralgias &/or arthritis—often the knees &/or ankles, but other joints can be affected. May be the first sign of Behçets. X ray is normal but synovium often has high neutrophils or mononuclear cells & a vasculitis process. 43	Arthralgias, fatigue and malaise. No permanent joint or bone issues noted, and patients are symptom-free between PFAPA flares. 40,41	Arthralgias may come before the arthritis. 88% have polyarticular or oligoarticular arthritis, most often in the wrists, knees, &/or ankles. Some w/cervical spine, hip, temporomandibular joint arthritis or synovial cysts. 45,46	intercarpal and carpo- metacarpal joint space narrowing a few yrs. after onset of AOSD–25%	
VASCULITIS	Not noted. <sup>1</sup>	Not noted. <sup>1</sup>	Vasculitis rarely develops. <sup>1</sup>	Vasculitis noted in 20% of patients. <sup>13</sup>	nodosa.1	HSP, lymphocytic vasculitis. <sup>1</sup>	Cutaneous vasculitis common, HSP is rare.	Not noted. <sup>11</sup>	A few w/localized or cerebral vasculitis. <sup>16</sup>	Not noted. <sup>18,53</sup>	Some w/Takayasu arteri- tis, or ANCA+ Vasculitis. <sup>54</sup>		Not noted. <sup>29</sup>	Some w/vasculitis, leuko cytoclastic vasculitis. <sup>34</sup>		Not noted. <sup>26,27</sup>	Extensive vasculits. 30% w/venous thrombosis. 43		Not noted. <sup>45,46</sup>	Not noted. <sup>44</sup>	Not noted. <sup>47,48,49</sup>
AMYLOIDOSIS	amyloidosis in some patients. <sup>1,9</sup>	Elevated SAA >25 % w/secondary amyloidosis. <sup>1,9</sup>	Elevated SAA. Second- ary amyloidosis in <2% pts. <sup>1,6</sup>	A few patients have developed secondary amyloidosis. <sup>13</sup>	Common >50% in untreated patients, it depends on genotype. <sup>9</sup>		<5-10%—uncommon. <sup>9</sup>	Not noted-unknown. <sup>9,11</sup>	Not noted. 15,16,17	Not noted. <sup>18,53</sup>	Not noted. <sup>19,22,54</sup>	Not noted. <sup>23, 24</sup>	Not noted. <sup>29</sup>	Not noted. <sup>34</sup>	Not noted. <sup>39</sup>	Not noted. <sup>26,27</sup>	Not noted. <sup>42,43</sup>	Not noted. <sup>40,41</sup>	Amyloidosis occurs in 7.4% of pts. in the USA, and 16% in Turkey.46	Very rare. <sup>44</sup>	Not noted. <sup>47,48,49</sup>
ABNORMAL LABS	High: ESR, CRP, SAA. Leukocytosis with flares. <sup>1</sup>	High: ESR, CRP, SAA. Leukocytosis,with flares. <sup>1</sup>	Chronically high: ESR, CRP, SAA, anemia, granulocyte leukocytosis. <sup>1,6</sup>	Monoclonal IgM &/or IgG gammopathy. High: ESR, CRP. Leukocytosis. Complement normal to elevated. 50% w/inflam- matory anemia. <sup>13</sup>	Fibrinogen, Leukocyto-		High: ESR, CRP, SAA w/ flares. High IgD w/IgA in 80% pts. Mevalonate aciduria noted during flares. <sup>1</sup>	thrombocytopenia. High: ESR, CRP, SAA, CK, IgD,	High: ESR, CRP, leukocy- tosis, chronic anemia. 16,56	Congenital dyserythro- poietic anemia (CDA). High ESR. WBC can be normal, or elevated— neutropenia in infancy. Cultures negative. <sup>18</sup>		Mildly elevated WBC, CRP & ESR rarely elevated—only during flares of symptoms. <sup>56</sup>	Cultures of bone & skin are negative. Purulent synovial fluid full of neu- trophils. High w/flares: CRP, ESR, WBC. <sup>29,30,32</sup>	High CRP & ESR, ACE, immunoglobulins. Anemia, leukopenia, eosinophilia, hematuria, proteinuria, pyuria, abnormal LFTs (LF).34,36	Elevated CRP during flares. <sup>39</sup>	ESR, triglycerides. Some	Leukocytosis common. Normal–rarely elevated ESR or CRP. Some cryo- globulinemia, elevated factor VIII, fibrinolysis. <sup>43</sup>	High: ESR, CRP, WBC during flares–normal levels when not flar- ing. <sup>40,41</sup>	High: ESR, CRP, WBC, SAA, ferritin, aldolase. Elevated LFT's. Leuko- cytosis, thrombocytosis Anemia. 45,46	High: ESR, CRP, LFTs, ferritin. Low glycosy- lated ferritin. Leukocy- tosis, anemia common w/flares. Prolonged PTT (DIC risk.) <sup>44</sup>	High: ESR, CRP, triglycerides, LFTs, soluble CD25, ferritin. Low: platelets, fibrinogen. Low NK cell cytotoxic function, neutropenia, anemia. <sup>49</sup>
				Gy			Gri				Grj				No image available		nttp://semulanst			CRI	No image available
	· · · · · · · · · · · · · · · · · · ·	The CAPS rash is often more pronounced during flares. (NOMID Alliance pt. image)	, o.		. ma around the ankle. (Arthriti	TRAPS rash on the chest of a child. (Swiss Med Wkly. 2012;142:w13602)		MA: 21 month old pt. w/ facial dysmorphism. (Orphanet Journal of Rare Diseases 2006 1:13)	SIS. (N Engl J Med 2009; 360:2426-		1'	CAMPS. (dermatlas.med.jhmi.edu,	/ PAPA: Pyoderma gangreno- sum. (dermatlas.med.jhmi.edu/image/ pyoderma_gangrenosum_1_020918)	brown papules. (Dermatology	Please contact nomidalli- ance.org if you can share an image of NLRP12.	1 ' ' '	1 3	PFAPA: Aphthous stomatitis (drpaulose.com; blog.timesunion.com/ mdtobe Mystery Monday 125)	on abdomen. (Joint Bone Spine,		Please contact nomidalli- ance.org if you can share an image of Familial HLH.

Main authors: Karen Durrant RN, BSN-President of The NOMID Alliance & Dr Juan Ignacio Aróstegui MD-Department of Immunology: Hospital Clínic de Barcelona. Acknowledgements: A special thanks to many medical doctors who have helped to make voluntary suggestions in regards to this reference chart: Dr Juan Ignacio Aróstegui, Dr Hal Hoffman, Dr Raphaela Goldbach-Mansky, Dr Anna Simon, Dr Polly Ferguson, Dr Rebecca Marsh, Dr Daniel Kastner, Dr Luca Cantarini, & Dr Lori Broderick. Thank you Nathan Durrant for donating your graphic design services. We want to thank all the doctors from the International Society of Systemic Auto-Inflammatory Diseases (ISSAID) for their research and dedication to patients with autoinflammatory diseases, and for the opportunity to present this chart at the Autoinflammation 2013 Congress. Our deepest thanks to The NOMID Alliance Board of Directors and to all the patients who have supplied images for this chart, & support for The NOMID Alliance. You are our greatest inspiration and strength!

Disclosure: All the individuals involved in the authorship, review, editing and creation of this chart voluntarily donated their help for this educational reference, and received no financial compensation. Permission granted by The NOMID Alliance to have this poster printed and distributed in the UK, The Republic of Ireland, The Netherlands, Belgium and Luxembourg by Swedish Orphan Biovitrum Ltd. for educational purposes only.

## **List of abbreviations:**

ACE: Angiotensin-converting enzyme (lab test) ANCA+ Vasculitis: Granulomatosis w/polyangiitis (GPA); Wegener's GI: Gastrointestinal (organs in the abdomen) ARDS: Acute Respiratory Distress Syndrome CD25: Soluble interleukin-2-receptor CNS: Central Nervous System (involving the brain, spinal cord) CRP: C-reactive protein (lab test); DIC: Disseminated intravascular coagulation

**ESR:** Erythrocyte sedimentation rate (lab test); Westergren ESR HSP: Henoch–Schönlein purpura, anaphylactoid purpura ICP: Intracranial pressure LFTs: Liver function tests (lab test): AST, ALT, GGT, ALK Phos, Bilirubin w/: abbreviation for the word "with" NK cells: Natural killer cells PMNs: Polymorphonuclear leukocytes (on lab tests w/ WBC count)

pt.: abbreviation for the word "patient" PTT: Partial thromboplastin time (lab test) SAA: Serum amyloid A protein (lab test) **TSH:** Thyroid-stimulating hormone (lab test); thyrotropin WBC: White Blood Count (lab test)

## All Cited References & Image Credits are Listed on the Back Side of this Chart.

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