SCHNITZLER'S SYNDROME, AN UNDERDIAGNOSED AUTOINFLAMMATORY DISEASE: CURRENT AND FUTURE PERSPECTIVE

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Abstract

Schnitzler's syndrome is a rare but usually underdiagnosed autoinflammatory disease, characterized by monoclonal IgM gammopathy, persistent urticaria, intermittent fever, bone pain, and arthralgia or arthritis. Although first reported by Dr. Liliane Schnitzler in 1972, the syndrome continues to pose challenges in diagnosis, and it usually takes more than five years before it is properly identified. Disorders show up in a form of urticaria, arthritis, organomegaly, fever, lymphadenopathy, high ESR, leukocytosis, and bone pain. Central to its pathogenesis are immunologic perturbations and activation of the inflammasome. It is typically diagnosed through clinical exam with history and important feature identification such as monoclonal gammopathy and acute or relapsing urticarial rash. Therapeutically synthetic agents include anakinra, canakinumab, rilonacept, and anti-IL-6 have been used with variable success. Recent research also demonstrated how natural treatments such as Terminalia chebula, Emblica offcinalis, Schinus terebinthifolia, tulsi, asafoetida, and Wedelia plants have the potential for controlling the symptoms and changing inflammatory paths. Promising medicinal possibilities for these plants, according to in-silico investigations, point to further research into their clinical uses. With new therapy paths that promise better patient outcomes, Schnitzler's illness is overall a therapeutic challenge.

Keywords: Schnitzler's syndrome, auto-inflammatory, Wedelolactone, IgM or IgG, monoclonal antibody, NLRP3 mosaicism, NLRP3 inflammasome.

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1 1 Introduction

Recurrent fever rash, bone and joint pain, swollen lymph nodes, exhaustion, 2 a monoclonal IgM component, leukocytosis, and systemic inflammatory response 3 are indicative of the uncommon and underdiagnosed adult disease known as 4 Schnitzler's syndrome. Schnitzler syndrome is an unusual but treatable disease 5 distinguished by persistent urticaria, intermittent fever, bone pain, arthralgia or 6 arthritis, and monoclonal IgM gammopathy. Dr. Liliane Schnitzler, a French 7 dermatologist, characterised it first in 1972, and more than 40 patients have 8 subsequently reported it. In most cases, the disease goes undiagnosed for more than 9 5 years. This syndrome's wide range of clinical symptoms involves dermatologists, 10 internists, rheumatologists, and haematologists. This has become an important 11 concern because major morbidities have been experienced by patients with systemic 12 inflammation when left untreated. It has been considered an adult-onset auto-13 inflammatory disease [1-4]. 14

15

SYMPTOMS

A number of symptoms are associated with Schnitzler's syndrome including urticaria, arthritis, organomegaly, fever, lymph nodes, enlarged liver or spleen, high ESR, leukocytosis, bone pain, and anomalies in haemopoiesis [5] and more symptoms involves like increased platelet count due to thrombocytosis, neutrophil leukocytosis, monoclonal gammopathy, increasing ESR or CRP, decreasing complement levels, anaemia associated with a chronic disease, and increased IgM levels [6].

23

AETIOLOGY

Aside from its unknown exact origin, immune system dysregulation is the main cause of Schnitzler's syndrome, an auto-inflammatory disease characterized by high inflammatory markers, joint pain, fever, and recurring urticarial rashes. The most salient feature is the monoclonal gammopathy of IgM or IgG. [7-8].

28

PATHOPHYSIOLOGY

Immune system dysregulation, inappropriate inflammasome activation, and overproduction of proinflammatory cytokines are the hallmarks of Schnitzler syndrome, an autoinflammatory disorder [9]. The cause of chronic inflammation and increased inflammasome activation could be due to NLRP3 inflammasome mutations induced by somatic NLRP3 mosaicism (Figure 1) [10-11].

34 **DIAGNOSIS**

Clinical diagnosis and diagnosis by patient history are essential for the 35 diagnosis of Schnitzler's syndrome. The most critical diagnostic findings include 36 monoclonal gammopathy, usually IgMk, sometimes IgG, and urticarial rash resistant 37 to antihistamines, and persistent or recurrent. The presence of a monoclonal 38 gammopathy, specifically IgMk, in conjunction with clinical signs such as urticaria 39 and systemic inflammation, is usually sufficient to make the diagnosis of SchS. 40 Examples of SchS patients who do not exhibit gammopathy but respond favourably 41 to IL-1 treatment suggest that the diagnostic criteria may be overly stringent. Even 42 in the absence of gammopathy, laboratory abnormalities such as high ESR, CRP, 43

and particular skin biopsy results can help in the diagnosis of Schnitzler-like 44 disorders [12-15]. 45 Erythrocyte Sedimentation Rate (ESR): A slightly elevated level, a 46 sign of inflammation. 47 C-Reactive Protein (CRP): Another indicator of inflammation that is 48 elevated. 49 Interleukin-6 (IL-6) and IL-2 Levels: Elevated, suggesting the 50 presence of an inflammatory condition. 51 Monoclonal IgM Component Serum protein electrophoresis revealed 52 it, which is a crucial characteristic of Schnitzler's syndrome. 53 Normal TNF-α and IL-8 Levels: These offer more context but do not 54 rule out Schnitzler's syndrome. 55 Normal Creatinine, Complete Blood Count (CBC), and Serum 56 **Calcium**: These typical outcomes aid in ruling out further diseases. 57 No Bence-Jones Proteinuria: This aids in ruling out multiple 58 myeloma. 59 Normal Skeletal X-rays, Bone Marrow Aspirates, and Histology: 60 These results aid in excluding further hematologic and skeletal diseases. 61 Normal Cerebral and Thoracic Scans, Abdominal Sonography: The 62 normal imaging studies aid in excluding further systemic illnesses. 63 **TYPES OF TREATMENT** 64 Different treatment modalities exist for the comprehensive management of 65 Schnitzler's Syndrome, which altogether can be categorized as synthetic and natural 66 ways, each of which gives a unique role in symptomatic relief and treatment of the 67 mechanisms underlying the autoinflammatory disorder. 68 SYNTHETIC TREATMENT FOR SCHNITZLER'S SYNDROME 69 Before these treatments, the syndrome could not be effectively treated with 70 more than thirty drugs. Although substantial dosages of corticosteroids were 71 necessary for some patients, antihistamine treatment had no impact. Low serum 72 amyloid is the goal of IL-1 blocking treatments like rilonacept, canakinumab, and 73 74 anakinra [16]. 1. Anakinra 75 Recombinant IL-1RA anakinra binds to the IL-1 receptor very firmly but does 76 not cause signal transduction. It was initially licenced to treat rheumatoid arthritis, 77 and then it was also approved to treat deficiency of IL-1 receptor antagonists and 78 neonatal-onset multisystem inflammatory disease. Martinez-Toboada et al. 79 documented the first instance of anakinra being successfully treated to Schnitzler's 80 syndrome, with a high effectiveness rate of up to 94%. Nonetheless, certain studies 81 propose utilising the anakinra therapy response as a standard for Schnitzler's 82 syndrome. Anakinra is highly effective in treating Schnitzler syndrome; 83% of 83 patients experienced complete remission and 17% experienced partial remission, 84 according to observational research. Another group of 21 Schnitzler syndrome 85 patients experienced both full and partial remission, and all of them saw long-term 86

benefits with anakinra treatment [17-22]. 87

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Canakinumab 2.

Human monoclonal antibody canakinumab is used to treat a variety of 89 systemic juvenile idiopathic conditions. such arthritis, FMF. as 90 hyperimmunoglobulin D syndrome/mevalonate kinase deficiency, MWS, TRAPS, 91 older familial cold auto-inflammatory syndrome, and CAPS in both adults and 92 children. 34 Schnitzler syndrome patients received treatment in all, and 58.6% of 93 them reported having a full recovery. Over nine months, eight patients had a 94 persistent therapeutic benefit. In a randomised, placebo-controlled research, 95 canakinumab was used to treat five out of seven individuals, while no response was 96 seen in the 13 patients who received a placebo. All patients demonstrated a durable 97 response to canakinumab over 4 years, with normalising levels of SAA and C-98 reactive protein. This demonstrates canakinumab's ongoing treatment efficacy for 99 Schnitzler's syndrome [23-26]. 100

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88

Rilonacept 3.

A recombinant fusion protein called rilonacept targets IL-1 receptors and 102 inhibits them from attaching to the IL-1 receptor found on cell membranes. The 103 European Medicines Agency and the US Food and Drug Administration have 104 authorised it for the treatment of CAPS. Eight individuals in a clinical trial who 105 satisfied Lipsker diagnostic criteria were given a loading dosage of 320 mg and then 106 a 160 mg subcutaneous injection per week. GPA and the Schnitzler activity score 107 were used to gauge the clinical results. Three patients had a partial response, and 108 fifty per cent of patients had complete or almost complete remission at the one-year 109 follow-up. Within 24 hours of the initial dosage, the therapeutic effects started to 110 take effect [27-31]. 111

112

Anti-IL-6 treatment 4.

Treatment outcomes for Schnitzler's syndrome with tocilizumab, an anti-IL-6 113 drug, have been inconsistent. When tocilizumab was infused, some patients went 114 into complete remission, while others did not react well. In one clinical research, 115 tocilizumab helped some patients at first, but with time, those benefits were lost. It 116 was also mentioned that tocilizumab medication may not have been as beneficial for 117 certain individuals in Japan. In general, individuals with Schnitzler syndrome who 118 do not respond to IL-1 blockers may be candidates for tocilizumab monotherapy; 119 however, for improved results, it may also be administered in combination with other 120 immunosuppressants. To learn more about the long-term effectiveness and possible 121 recurrence rates of tocilizumab treatment for Schnitzler's syndrome, more 122 investigation and observation are required [32-36]. 123

124

Rituximab 5.

Rituximab is a monoclonal antibody that is used to treat ANCA-associated 125 vasculitis, rheumatoid arthritis, and B-cell lymphoma. Although no clinical trials 126 have been done, it might be useful in treating Schnitzler's syndrome. When used in 127 conjunction with other medications, rituximab has demonstrated a good response in 128 certain cases but not in others [37-44]. 129 6.

130

Anti-TNF

7.

Etanercept, adalimumab, and drome are examples of TNF inhibitors. Only one article demonstrates that adalimumab There was no improvement or escalation of symptoms seen [45,46].

134

Anti-IL-17 therapies

Schnitzler syndrome patients have lesional skin that contains IL-17A, and neutrophils are the source of this substance. Being a potent neutrophil chemoattractant, IL-17A causes cutaneous lesions and dermal infiltration. Those who do not respond to IL-1 blockers may benefit from anti-IL-17 therapy [45-51].

139

NATURAL TREATMENT FOR SCHNITZLER SYNDROME

Schnitzler Syndrome can be effectively treated with medicinal plants, the
formulation of rare autoinflammatory conditions. Table 1 presents traditional plants
used for preventing and treating this condition.

143

1. Terminalia Chebula

The medicinal herb Terminalia chebula, which has a wide range of therapeutic 144 uses, is traditionally used to treat urticaria, arthritis, fever, organomegaly, and 145 lymphadenopathy [52]. Furthermore, studies have demonstrated the potential of 146 Terminalia chebula extract to ameliorate doxorubicin-induced haematological 147 problems by enhancing haemoglobin content, total white blood cell count, and 148 antioxidant status [53]. Furthermore, research on Terminalia chebula has 149 demonstrated its antioxidant, anti-inflammatory, and anticancer properties, 150 indicating that it is a suitable option for additional study in cancer treatment [54– 151 55]. 152

The wide array of the pharmacological actions of Terminalia chebula might, 153 therefore, be helpful in some ways while treating the various manifestations of 154 155 Schnitzler's syndrome, although the contexts cited do not relate to its efficacy in any specific symptom pertinent in the case, such as elevated ESR, leukocytosis, and 156 abnormalities of hemopoiesis. Chelagic acid isolated from Terminalia chebula 157 reduces inflammation through the modification of proinflammatory cytokines. 158 Studies have shown that chebulagic acid inhibited the increase in IL-1 β and TNF- α 159 production induced by LPS. It has also been proven to inhibit the production of pro-160 inflammatory cytokines like TNF- α , IL-6, and IL-1 β through repression of the p38 161 MAPK pathway. It was also established that chebulinic acid exhibited anti-162 inflammatory activity by the inhibition of protein denaturation, which may justify 163 further study of the anti-inflammatory properties of the plant. Together, these results 164 point to the importance of chebulagic acid in modulating proinflammatory cytokines 165 by enhancing osteoblast functions, decreasing oxidative stress, and inhibiting NF-166 κ B signalling, as demonstrated in studies on LPS-induced bone loss. This provides 167 a promising therapeutic approach for inflammatory conditions such as Schnitzler 168 syndrome. [56,57]. 169

170

In Silico studies of Terminalia Chebula

The docking studies of Terminalia chebula unravel a great therapeutic
 potential due to its good binding affinity with the amino acids as shown in Figure 2.
 The phyto-constituents of T. chebula are one day going to be developed into potent
 medication options.

• The molecular dynamics simulations show the stability of these interactions and prove their potential for medication development. Experiments in vitro confirmed the antioxidant activities of T. chebula, showing that it can block oxidative reactions, a critical function.

• Even though the docking experiments are very promising for the species, further in vivo studies will be necessary to establish its therapeutic efficacy and safety profile for clinical application.

182

2. Emblica Officinalis

Ellagic acid, a substance with strong anti-inflammatory qualities, is found in 183 Emblica officinalis, commonly referred to as Indian gooseberry or Amla. Because 184 ellagic acid has been shown to target proinflammatory cytokines, it may be able to 185 treat inflammatory disorders such as Schnitzler syndrome. This has been the subject 186 of much research. According to studies, ellagic acid's anti-inflammatory properties 187 are achieved through modulating inflammatory signals and cytokines, including 188 TNFα, IL-1β, and IL-6. AKT1, VEGFA, TNF, and MAPK3 are some of the 189 important targets of ellagic acid that have been linked to inflammation, according to 190 research employing a network pharmacology approach. It has been demonstrated 191 that ellagic acid administration decreases pro-inflammatory cytokine levels, such as 192 TNF- α and IL-1 β , while raising anti-inflammatory cytokines, such as IL-10 and 193 IFN-y, hence lowering arthritis-associated pathology in animal modelsThese 194 findings support the fact that ellagic acid contained in Emblica officinalis could 195 modulate the effects of Schnitzler's syndrome, by its action on inflammation 196 responses through various different molecular pathways and focusing particularly 197 on proinflammatory cytokines [58-61]. 198

199

In Silico studies of Emblica Officinalis

• Emblica officinalis has a good binding affinity with amino acids, and its complex phytochemical profile makes it a potential option for medicinal application as seen in Figure 2. This action will not only make it highly bioactive but also place it in an excellent position for use in modern medicine.

• Its ability to complexation with amino acids means that the absorption and, therefore, the efficacy of these medicaments would be higher and therefore better outcomes for patients.

• While most of the medicinal potential for Amla has already been well documented, further studies are needed to understand the mechanism of its interaction with amino acids and to standardize its extracts for clinical use.

210

3.

Schinus Terebinthifolia

Traditional medicine used Schinus terebinthifolia, sometimes referred to as 211 capoeira, to treat a range of ailments, including inflammatory diseases like arthritis 212 [62]. The herb exhibits anti-inflammatory and restorative qualities, but there isn't 213 much research on how it affects certain conditions like Schnitzler syndrome. 214 Monoterpenes, sesquiterpenes, and phenols are among the bioactive components 215 found in Schinus terebinthifolia, and they may be beneficial for illnesses involving 216 organomegaly, fever, lymph nodes, and hemopoiesis dysregulation [63]. With the 217 low cytotoxicity and antibacterial activity, it becomes equally important to 218

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understand the therapeutic benefits and limitations of the essential oil from Schinus 219 terebinthifolia against a wide range of health conditions [64]. Gallic acid is one such 220 polyphenolic component of Schinus terebinthifolia and many other natural sources, 221 and there are various ways it can affect the levels of proinflammatory cytokines. 222 TNF- α , IL-1 β , and IL-6 are the proinflammatory cytokines which gallic acid 223 effectively reduces by downregulating its gene and inhibiting the signaling pathways 224 such as NF-κB [65–67]. It also exerts anti-inflammation by inhibition of myometrial 225 contractions, reducing the expression of labour-associated proteins in fetal tissues, 226 and regulating the production of proinflammatory mediators and cytokines [68]. 227 Further, gallic acid has shown anti-inflammatory and antioxidant action in respect 228 of polycystic ovarian syndrome by reducing the amount of oxidative stress, 229 inflammatory cytokines, lipid peroxidation, and DNA oxidative damage in ovarian 230 tissue [69]. All things considered, gallic acid is a viable option for the treatment of 231 inflammatory illnesses and disorders due to its capacity to regulate proinflammatory 232 cytokines. 233

234

In-Silico's studies of Schinus Terebinthifolia

In-silico studies on Schinus terebinthifolia (Fig. 2), focusing on gallic acid, 235 have remarked on its remarkable binding affinity with multiple amino acids, 236 showing much potential as a therapeutic agent. In-silico studies use the molecular 237 docking and computing study interactions of gallic acid against various target 238 proteins. Such studies suggest that gallic acid and other phytochemicals could be 239 encouraging candidates for drug development in thrombin activity and skin 240 disorders. Such findings are in need of further validation in vitro and in vivo to 241 confirm the findings and find out the fullest therapeutic potentiality of Schinus 242 terebinthifolia. 243

244

4. Tulsi Plant

Tulsi, scientifically known as Ocimum sanctum, is a highly valued medicinal 245 herb with diverse therapeutic uses [70–73]. Traditionally, this plant has been used in 246 Ayurveda and other forms of traditional medicine for the management of various 247 diseases and disorders, including but not limited to organomegaly, urticaria, arthritis, 248 fever, disorders of lymph nodes, and abnormalities of haemopoiesis. Some of the 249 major medicinal values of tulsi are attributed to its anti-inflammatory, antioxidant, 250 and immunomodulatory properties. These uses are facilitated by the phytochemical 251 elements found in different portions of the plant, including as eugenol, vallinin, 252 gallic acid, and linoleic acid [70,71,73]. These qualities make tulsi a useful natural 253 therapy for treating a range of health issues, possibly including those related to 254 Schnitzler's syndrome [72], while particular clinical research on this illness has not 255 yet been done. The plant tulsi (Ocimum sanctum) and other plants contain ursolic 256 acid, a pentacyclic triterpenoid that has been shown to have powerful anti-257 inflammatory effects. Important inflammatory cytokines like IL-6, IL-8, TNF-a, IL-258 1β, and IL-18 can be inhibited by ursolic acid, according to studies. Many disorders, 259 such as psoriasis, cerebral haemorrhage, and chronic prostatitis, have been 260 discovered to have their inflammatory responses attenuated by ursolic acid through 261 numerous routes, including the NF-kB/NLRP3/GSDMD pathway [74–77]. 262

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263 264

In-Silico's studies of Tulsi

The in-silico studies regarding Tulsi (Ocimum sanctum) were conducted with molecular docking and analysis of bioactive components that indeed 265 pointed out the plant's great binding affinity with several amino acids as portrayed 266 in Figure 2. In our findings, draws attention to the significant role Tulsi plays in the 267 treatment of a number of medical disorders within the purview of modern medicine, 268 and now it is becoming critical in drug discovery and development. 269

270 271

It has been reported that tulsi has a wide range of pharmacological action, including anti-bacterial, anti-inflammatory, and anti-cancer activities.

272

5. Asafoetida

Asafoetida, which is derived from the Ferula asafetida plant, has been used 273 traditionally to treat several diseases that are linked to Schnitzler's syndrome, 274 including urticaria, arthritis, fever, organomegaly, lymph nodes, enlarged liver or 275 spleen, elevated ESR, leukocytosis, bone soreness, and irregularities in hemopoiesis 276 [78-82]. The plant is good for a variety of diseases due to its therapeutic 277 characteristics, which are attributed to its bioactive components, which include 278 coumarins, volatile oils, and ferulic acid. Antifungal, anti-inflammatory, antiviral, 279 and antioxidant activities make asafoetida useful in food preparation owing to its 280 immense pharmacological importance. Galvanic acid is among the identified 281 compounds in asafoetida, which indicates the anti-cancerous activities of the plant, 282 further indicating various medicinal applications of the plant. Anti-inflammatory 283 activity has been reported with Ferula assa-foetida-an asafoetida supply plant-by 284 modulation of proinflammatory cytokines especially in its ethanolic extract fraction. 285 It is due to studies that have shown the ethanolic extract of Ferula assafoetida oleo-286 gum-resin significantly reduced intracellular ROS formation and suppressed the 287 expression of ICAM-1 and VCAM-1; there is less adhesion of PBMCs [83]. The 288 anti-inflammatory action of Ferula assa-foetida gum-resin extract has been attributed 289 to its rich content of antioxidant flavonoids and phenolic components which inhibit 290 lipoxygenase activity [84]. Additionally, it has been documented that member of 291 the genus Ferula, especially Ferula assa-foetida, influence different cytokines and 292 reduce inflammatory mediators, demonstrating their potential as therapeutic agents 293 against inflammatory diseases [85]. 294

295

In-Silico's studies of Asafoetida

The interactions of ferulic acid with amino acids at the active site of 296 proteins, especially Ferula asafoetida, offer a pharmacological potential for it. In-297 silico studies revealed that ferulic acid binds well with the key amino acids to 298 enhance bio-activity and point toward the therapeutic application as shown in Figure 299 2. 300

301 • However, while these computational findings turn out to be promising, further experimental validation needs to be done to confirm the very mechanism of 302 binding and understand the full therapeutic potential of Asafoetida. 303

304

Wedelia Plant

6.

Many studies have been conducted on the potential medical benefits of 305 wedelia plants, especially Wedelia chinensis, for treating a variety of conditions, 306

including fever, coughing, phlegm, and liver damage [86-88]. Wedelia chinensis 307 has been reported to contain chemicals with important biological activities, such as 308 steroids and diterpenes, which have been proven to have anti-acetylcholinesterase, 309 antioxidant, and cytotoxic properties [86]. Some of these substances may be 310 medically useful. Antifungal activities of Wedelia species were also investigated and 311 some interesting compounds were isolated that showed promising potentials in 312 further studies and development into medication [89]. Induction of polyploidy in 313 Wedelia chinensis was also conducted to increase the quality and medicinal value of 314 the plant, therefore demonstrating the usefulness and potential therapeutic uses of 315 the plant [88]. Accordingly, the great variety of bioactive compounds and medicinal 316 properties within the diverse species of Wedelia would show a very wide range of 317 potential health benefits that could be investigated against a number of conditions, 318 including those related to Schnitzler's syndrome-even though these contexts do not 319 specifically mention the connection between Schnitzler's syndrome and Wedelia 320 plants. Wedelolactone is obtained from a wedelia plant and has anti-inflammatory 321 properties due to the ability to modulate pro-inflammatory cytokines through various 322 In collagen-induced arthritis (CIA) mice, studies reveal that pathways. 323 wedelolactone reduces the release of pro-inflammatory cytokines like IL-1β, IL-6, 324 TNF- α , and IL-18 [90]. Furthermore, Wedelolactone has been observed to potentiate 325 PKA signalling to extensively decrease NLRP3 inflammasome activation, 326 pyroptosis, and IL-1ß secretion in macrophages [91]. Furthermore, in murine 327 macrophages driven to an inflammatory response by zymosan, wedelolactone 328 inhibits the release of TNF-a, IL-6, and IL12p40 [92]. Additionally, Wedelolactone 329 inhibits oxidative stress and inflammation by targeting soluble epoxide hydrolase 330 (sEH) to increase EET levels and regulate Nrf2 and NF-kB pathways in LPS-331 stimulated acute lung damage models [93]. Together, our results highlight the 332 anti-inflammatory characteristics of wedelolactone in diverse regulating 333 proinflammatory cytokines via several mechanisms. 334

335

In-Silico's studies of Wedelia Plant

Wedelolactone, the bioactive compound extracted from Wedelia plant, 336 showed prominent binding interactions with amino acids as shown in Figure 2, 337 proving the potential of the compound as a drug. The phytoconstituents present in 338 this plant are amino acids, flavonoids, and phenolic compounds, enhancing its 339 nutritional and therapeutic value. Wedelia trilobata has potent anti-inflammatory 340 activities due to its interaction with amino acids. The potential of Wedelolactone for 341 the treatment of diseases is varied, thus justifying the possibility of its positive 342 interaction with amino acids. Further research is essential to clearly understand the 343 mechanism behind these interactions and their therapeutic applications. 344

345 2 Conclusion

Characterized by chronic urticaria, recurring fever, bone pain, arthralgia or arthritis and monoclonal IgM gammopathy, Schnitzler's syndrome is a rare but treatable disorder in adults. It is a rare auto-inflammatory disease that often remains unrecognized for more than five years. Symptoms include urticaria, arthritis, fever, lymph nodes, swollen liver, or spleen, raised ESR, leukocytosis, bone pain, and

abnormalities in haemopoiesis. Immune system dysregulation is the major cause of 351 352 this autoinflammatory condition. Synthetic treatment for Schnitzler's syndrome includes anakinra, canakinumab, 353 rilonacept, and a combination of these drugs. Anakinra is highly effective in treating 354 Schnitzler syndrome, with 83% of patients experiencing complete remission and 355 17% experiencing partial remission. Canakinumab is used to treat various 356 juvenile idiopathic arthritis, conditions. including systemic FMF. 357 hyperimmunoglobulin D syndrome/mevalonate kinase deficiency, MWS, TRAPS, 358 older familial cold auto-inflammatory syndrome, and CAPS in both adults and 359 children. Medicinal plants have shown potential benefits in managing some aspects 360 modifying proinflammatory cytokines, of the syndrome by inhibiting 361 lipopolysaccharide-induced elevation of interleukin (IL)-1ß and tumour necrosis 362 factor (TNF)- α , and enhancing osteoblast functions. Terminalia chebula, Emblica 363 Officinalis, Schinus Terebinthifolia, Tulsi Plant, Asafoetida, Wedelia plants, and 364 Wedelolactone have shown potential in treating Schnitzler's syndrome. In-silico 365 studies on Terminalia Chebula and Tulsi have highlighted its great binding affinity 366 with amino acids, highlighting its significant role in modern medicine and drug 367 discovery and development. Asafoetida has been traditionally used to treat various 368 diseases linked to Schnitzler's syndrome, and Wedelolactone has anti-inflammatory 369 properties due to its ability to modulate proinflammatory cytokines through various 370 routes. Further research is needed to understand the mechanism behind these 371 interactions and their therapeutic applications. 372

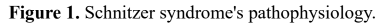
- 373 **Consent for Publication**
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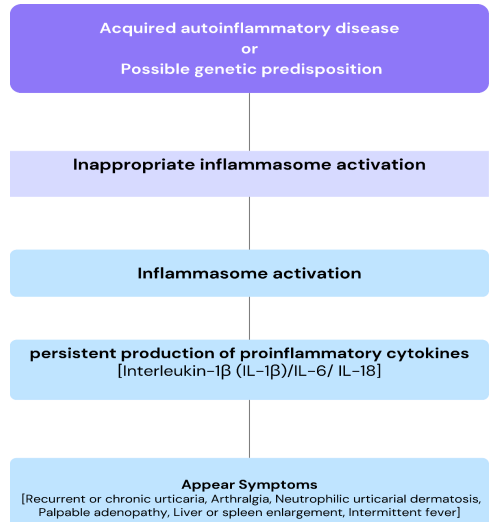
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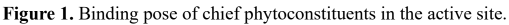
Table 1.	List	of	Medicinal	Plants	Used	for	the	Prevention	and	Treatment	of
Schnitzle	r's syr	ndro	ome.								

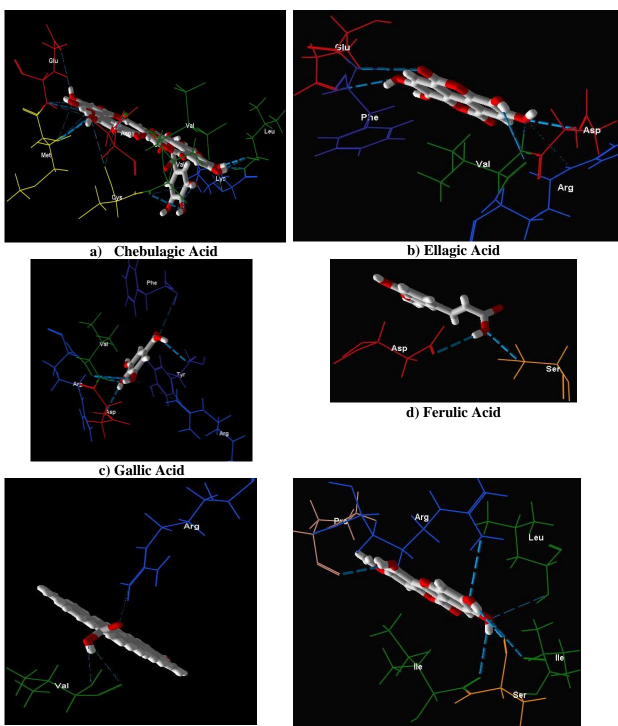
S.NO.	PLANT	FAMILY	CHEMICAL COMPOSITION USED IN SCHNITZLER'S SYNDROME	COMMON NAME	PART OF THE PLANT USED
1	Terminalia Chebula	Combretaceae	Chebulagic acid,	Black Myrobalan	Fruit
2	Emblica Officinalis	Phyllanthaceae	Ellagic acid	Amla	Fruit, Seed, Leaves, Root, Bark and Flowers
3	Schinus Terebinthifolia	Anacardiaceae	Gallic acid,	Brazilian Peppertree	Bark, leaves
4	Tulsi Plant	Lamiaceae	Ursolic acid	Holy basil	Leaves, stem, flower, root, seeds
5	Asafoetida	umbelliferae	Ferula assa- foetida	Devil's dung, Hing	Root, Leaves, and stem
6.	Wedelia Plant	Asteraceae	Wedelolactone	Yellow Dots	leaves

РИСУНКИ

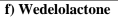








e) Ursolic Acid



ТИТУЛЬНЫЙ ЛИСТ_МЕТАДАННЫЕ

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Блок 3. Метаданные статьи SCHNITZLER'S SYNDROME, AN UNDERDIAGNOSED AUTOINFLAMMATORY DISEASE: CURRENT AND FUTURE PERSPECTIVE

Сокращенное название статьи для верхнего колонтитула:

Keywords: Schnitzler's syndrome, auto-inflammatory, Wedelolactone, IgM or IgG, monoclonal antibody, NLRP3 mosaicism, NLRP3 inflammasome.

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