

1 **TITLE**

2 Kynurenine monooxygenase blockade reduces endometriosis-like
3 lesions, improves visceral hyperalgesia, and rescues mice from a
4 negative behavioural phenotype in experimental endometriosis.

5

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31 **SUMMARY**

32 Endometriosis is a common and debilitating neuro-inflammatory disorder that is associated with chronic pain.
33 Definitive diagnosis is based on the presence of endometrial-like tissue (lesions) in sites outside the uterus.
34 Kynurenine monooxygenase (KMO) is a mitochondrial enzyme of tryptophan metabolism that regulates
35 inflammation and immunity. Here, we show that KMO is expressed in epithelial cells in human endometriosis
36 tissue lesions and in corresponding lesions in a mouse model of endometriosis. In mice, oral treatment with the
37 potent KMO inhibitor KNS898 induced a biochemical state of KMO blockade with accumulation of
38 kynurene, diversion to kynurenic acid and ablation of 3-hydroxykynurene production. In the mouse model
39 of endometriosis, KMO inhibition improved histological outcomes and endometriosis pain-like behaviours,
40 even when KNS898 treatment commenced one week after initiation of lesions. Taken together, these results
41 suggest that KMO blockade is a promising new non-hormonal therapeutic modality for endometriosis.

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43 143 words.

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45

46 **KEYWORDS**

47 Kynurenine monooxygenase; Endometriosis; Drug discovery; Non-hormonal therapies; Visceral hyperalgesia
48

49 INTRODUCTION

50 Endometriosis is a life-altering condition that affects approximately 10% of females. It is an oestrogen-
51 dependent neuroinflammatory disorder associated with debilitating pelvic pain, excessive fatigue,
52 gastrointestinal and urinary symptoms, and infertility¹. Worldwide, 200 million prevalent cases are forecast by
53 2026. Endometriosis is defined by the presence of endometrial-like tissue ('lesions') outside the uterus.
54 Physiological hormonal fluctuations in women induce cyclical episodes of cell proliferation, inflammation,
55 injury, and repair within lesions that favour fibroblast to myofibroblast differentiation and fibrosis¹. We and
56 others have identified metabolic dysfunction in cells associated with development of endometriosis lesions². At
57 present, therapeutic options are largely limited to surgery (that often needs to be repeated) or medical therapies
58 that target hormonal activity with resultant side effects (block conception, menopausal symptoms)¹. Patient
59 surveys consistently show frustration with the lack of available treatments that can give long term relief from
60 symptoms including pain, low mood and bloating³. Analysis of recent clinical trials directed at endometriosis¹
61 has highlighted an unmet need for new, non-hormonal approaches to symptom relief. The studies in the current
62 paper have addressed this need by focusing on an enzyme that is known to play a key role in inflammatory
63 processes that are implicated in the aetiology of endometriosis, but which has not previously been investigated
64 as a target.

65 Our proposed solution to this unmet medical need is by targeting the enzyme kynureine 3-
66 monooxygenase (KMO). KMO is a critical regulator of inflammation at multiple organ sites that acts by altering
67 metabolic flux through the kynureine pathway of tryptophan metabolism⁴. KMO is known to be expressed in
68 non-pathological endometrium⁵, but whether KMO is over-expressed in endometriosis lesions and linked to the
69 severity of inflammation remains to be determined. KMO has been identified as a critical step in converting
70 kynureine to the cytotoxic metabolite, 3-hydroxykynureine (3HK), that is an oxidative stressor, causes protein
71 cross-linking, and regulates the immune-metabolic interface⁴. Although there is no specific information about a
72 direct role of KMO in endometriosis, there is evidence of dysregulated tryptophan metabolism in a recent study
73 using a preclinical non-human primate model of endometriosis⁶, and increased kynureine pathway flux at the
74 immune-metabolic interface between stromal cells and NK immune cells in endometriosis lesions⁷.

75 At present, there is a scientific rationale for KMO inhibition in endometriosis, but it remains to be
76 shown in preclinical experiments whether KMO inhibition is efficacious in decreasing lesion volume or
77 behavioural symptoms which are used as a surrogate for pain responses in model systems. KNS898 is a highly
78 specific small molecule KMO inhibitor with potential for use by women with endometriosis, based on
79 favourable characteristics for oral development in terms of bioavailability and predicted half-life^{8,9}. KNS898 is a
80 competitive inhibitor of kynureine substrate at the active site of KMO with a pIC50 of 8.8⁸⁻¹⁰. We propose that
81 KMO inhibition is a novel therapeutic strategy for endometriosis and, if successful, we will make a significant
82 positive impact for women with this painful, disabling condition. The aim of this project was to obtain proof-of-
83 concept for KMO inhibition as a novel therapy for endometriosis. Specifically, we sought to explore the
84 expression of KMO in biobanked human endometrial and endometriosis lesion tissues, confirm target inhibition
85 of KMO by KNS898 in mice, and define the preclinical efficacy of KNS898 in improving clinical features of
86 disease (specifically hyperalgesia and altered cage behaviour) and reducing endometriosis lesion volume in an
87 experimental mouse model of endometriosis.

89 **RESULTS**

90 **KMO is expressed in human eutopic endometrium and human endometriosis tissue lesions.** To explore
91 whether we could detect variations between expression of KMO in endometrium (eutopic) within the uterus and
92 a variety of lesions obtained from patients, we conducted detailed immunohistochemistry with a primary
93 antibody specific for KMO. On fixed tissue sections of normal human endometrium KMO expression was most
94 striking in epithelial cells lining the glands (Figure 1a, insert B) with lower levels in the luminal layer (insert C).
95 Notably expression in the glands was not uniform (**Figure 1a**). KMO was also strongly immunopositive in
96 human peritoneal endometriosis lesions (**Fig. 1d to Fig. 1g**), and evidently mostly localised to the epithelial
97 tissues surrounding the distended endometrial gland-like structures (DEGLS) (**Fig. 1e and Fig. 1g**). Expression
98 in the stromal compartment appeared variable. In human ovarian endometriosis lesions, KMO was present at
99 low expression levels in the mesothelial layers (**Fig. 1h and Fig. 1i**). Duplex immunohistochemistry with cell
100 phenotype markers CD68 (macrophages) did not show KMO colocalising with these immune cells (data not
101 shown)

102

103 **Oral KNS898 inhibits KMO in mice.** Next, we established that oral dosing of KNS898 by gavage in mice
104 resulted in inhibition of KMO. Using n=3 mice per group, we administered KNS898 at 0.01 mg/kg, 5 mg/kg,
105 and 25 mg/kg twice daily (b.d.) in vehicle for seven days, as described. Plasma drug levels and metabolite
106 concentrations are shown in **Figure 2**. KNS898 dosed at 0.01 mg/kg b.d. resulted in a mean (\pm S.E.M.) plasma
107 drug level of 0.18 ± 0.01 ng/mL, 5 mg/kg resulted in 88.8 ± 22.6 μ g/mL, and 25 mg/kg gave 483.9 ± 84.0
108 μ g/mL. The difference between groups was statistically significant by one-way ANOVA with post hoc Tukey's
109 test ($P = 0.001$) (**Fig.2a**). KMO blockade with KNS898 was clearly measurable. A backlog in the KMO
110 substrate KYN was evident: KNS898 dosed at 0.01 mg/kg b.d. resulted in a mean (\pm S.E.M.) plasma level of
111 KYN of 339 ± 39 ng/mL, 5 mg/kg resulted in 4940 ± 483 ng/mL, and 25 mg/kg gave 3682 ± 634 ng/mL. The
112 difference between groups was statistically significant by one-way ANOVA with post hoc Tukey's test ($P =$
113 0.001). The increase in KYN at maximal inhibition was approximately 14-fold compared to the level seen after
114 KNS898 0.01 mg/kg (**Fig.2b**). Excess KYN was metabolised to KA by kynurenine aminotransferase: KNS898
115 dosed at 0.01 mg/kg b.d. resulted in a mean (\pm S.E.M.) plasma level of KA of 629 ± 103 ng/mL, 5 mg/kg
116 resulted in 14399 ± 3394 ng/mL, and 25 mg/kg gave 15965 ± 789 ng/mL. The difference between groups was
117 statistically significant by one-way ANOVA with post hoc Tukey's test ($P = 0.001$). The fold increase in KA at
118 maximal inhibition was approximately 25-fold compared to the level seen after KNS898 0.01 mg/kg (**Fig.2c**).
119 KMO blockade resulted in a statistically-significant reduction of 3HK in plasma: KNS898 dosed at 0.01 mg/kg
120 b.d. resulted in a mean (\pm S.E.M.) plasma level of 3HK of 27.7 ± 7.2 ng/mL, 5 mg/kg resulted in 4.2 ± 0.3
121 ng/mL, and 25 mg/kg gave 0.9 ± 0.4 ng/mL. The difference between groups was statistically significant by one-
122 way ANOVA with post hoc Tukey's test ($P = 0.001$). The fold decrease in 3HK at maximal inhibition was
123 approximately 30-fold compared to the level seen after KNS898 0.01 mg/kg (**Fig.2d**). Overall, there was a clear
124 dose response to KNS898 administration leading to maximal KMO blockade at 25 mg/kg b.d. This dose was
125 therefore selected for efficacy experiments going forward. A diagrammatic representation of the kynurenine
126 pathway is shown as Figure 2e for reference.

127

128 **KMO blockade reduces endometrial gland-like lesion burden in experimental endometriosis in mice.** The
129 experimental design for the mouse model of endometriosis is shown in **Figure 3a**. The pharmacological effect
130 of KNS898 therapy showed appropriate levels of KNS898 detected in plasma (**Fig 3b**), with accumulation of
131 kynurenone (**Fig. 3c**), inhibition of 3HK production (**Fig. 3d**) and diverted metabolism of accumulated
132 kynurenone to kynurenic acid (**Fig. 3e**). All recipient mice inoculated with donor tissue (groups G3, G4, and G5)
133 developed distended endometrial gland-like structures (DEGLS). The incidence of DEGLS formation was
134 enumerated at autopsy, and the axial length of each DEGLS was measured after excision from the surrounding
135 tissue. In G3 (endometriosis + vehicle), 8 of 15 (53%) of the inoculated animals had developed DEGLS. In
136 KNS898-treated group G4 (endometriosis + treatment from Day 19), DEGLS formed in 4 of 15 mice (26.7%)
137 and in G5 (Endo + treatment start on Day 26) in 6 of 15 mice (40%). As expected, no DEGLS were formed in
138 the non-inoculated control and sham groups. The total number of DEGLS per animal in each group was highest
139 in G3 with an average of 4.0 per animal with DEGLS (total = 32 DEGLS in 8 mice in G3). Mice with
140 endometriosis receiving KNS898 from the time of inoculation (G4) had an average of 2.0 DEGLS per animal
141 with DEGLS (total = 8 DEGLS in 4 mice in G4) and those receiving KNS898 1 week after inoculation (G5) had
142 an average of 1.8 DEGLS per animal (total = 11 DEGLS in 6 mice in G5) (**Figs. 3f and 3g**). Statistical analysis
143 by ANOVA showed a significant difference in endometriosis DEGLS burden between groups ($P = 0.0295$ for
144 DEGLS per animal; $P = 0.004$ for DEGLS per group). DEGLS axial length and derived volume did not differ
145 between groups (**Supplementary Fig. 1a and b**). All recipient mice inoculated with donor tissue lost body
146 weight following inoculation which then gradually recovered. After recovery, body weight of all three
147 inoculated groups was lower compared to the control groups for the duration of the study. Overall, there was no
148 significant difference in body weight between G3 and the KNS898 treatment groups G4 and G5 (endometriosis
149 + treatment from Day 26) (**Suppl. Fig. 1c**).
150

151 **KMO is expressed in experimental endometriosis in mice.** Histological examination of DEGLS identified
152 them as containing cystic structures lined with epithelial layers identifiable as columnar epithelium,
153 pseudostratified epithelium, squamous epithelium, and cuboidal epithelium, with goblet cells. These DEGLS
154 were considered to represent endometriosis-like lesions derived from the implanted basal
155 endometrial/myoepithelial layers of the donor mice uteri (**Suppl. Fig 2**). Immunohistochemistry using an
156 antibody to KMO showed KMO protein expression localised mainly to the epithelial cells lining of the DEGLS,
157 with a lesser degree of KMO positive staining in the closest surrounding connective tissue, in keeping with the
158 previously observed KMO expression pattern in human endometriosis lesion tissue (**Fig. 4a and 4d and**
159 **Supplementary Fig S3**). The thickness (area divided by length) of the KMO positive epithelial layer was
160 quantified for each DEGLS section using QuPath and there was no difference between groups G3, G4 and G5
161 (**Fig. 4b and 4e**). However, quantification of KMO expression confirmed the high intensity of KMO staining in
162 the epithelial lining layers (**Fig. 4c and 4f**), but also showed a clear and statistically significant reduction in
163 KMO expression intensity in those areas in DEGLS removed from mice treated with the KMO inhibitor
164 KNS898 (**Fig. 4g**; $P = 0.008$). Representative micrographs from each of groups G3, G4 and G5 are presented in
165 **Figures 4h, 4i and 4j**.
166

167 **KMO inhibition reduces mechanical allodynia in experimental endometriosis.** Clinical endometriosis is
168 associated with visceral hyperalgesia and central sensitisation to pain¹¹. Visceral and central hyperalgesia may
169 be tested in rodents using the Von Frey filament test¹². Baseline reaction values for hind paw and bladder Von
170 Frey tests showed no significant difference in mechanical allodynia before inoculation. In established
171 endometriosis without treatment (group G3), the mechanical allodynia threshold in the hind paw was
172 statistically significantly lower compared to baseline for the group. When compared to the control groups at the
173 corresponding time point beginning 1 week after inoculation and continuing until the end of the study. Day 26
174 KNS898-treated group (G5) showed a statistically-significant improvement in mechanical allodynia in the hind
175 paw using the Von Frey test compared to mice in G3 with untreated endometriosis given vehicle control (Two-
176 way ANOVA, Group effect P = 0.003, time effect P < 0.0001; Dunnett's multiple comparison test G5 vs G3
177 P=0.001) (**Fig. 5a**). The mechanical allodynia threshold for the bladder reflex also was lower in mice with
178 endometriosis compared to baseline throughout the study, and KNS898 treatment commencing at D26 (G5) was
179 associated with a statistically significant improvement in bladder mechanical allodynia threshold at D42
180 compared to mice with untreated endometriosis given vehicle control (G3)(Two-way ANOVA, Group effect P =
181 0.038, time effect P < 0.001; Dunnett's multiple comparison test G5 vs G3 P=0.021) (**Fig. 5b**).
182

183 **KMO inhibition rescues impaired cage exploration behaviour and mobility in mice with endometriosis.**
184 HCA peripheral moving speed, time at cage edge, and illness behaviour, including temperature, motility and
185 cage exploration behaviour was quantified using Home Cage Analysis (HCA). Baseline HCA was recorded
186 before inoculation and at the end of the experiment. Mice with endometriosis without treatment showed an
187 overall reduction in activity in moving distance and moving speed relative to baseline, and compared to sham-
188 operated control groups, indicating a negative effect on behaviour due to endometriosis. Importantly, mice with
189 endometriosis treated with KNS898 showed marked improvement in motility and cage exploration behaviour
190 compared to untreated endometriosis mice, and although this difference between groups was statistically
191 significant by Welch's one-way ANOVA, post hoc testing (Dunnet's T3) was not significant between groups.
192 This qualitative difference of time spent exploring the periphery of the cage being lower for mice with
193 endometriosis treated with vehicle control was seen in both the day and night phases (**Figs. 5c and 5d**). A
194 similar improvement with KNS898 treatment compared to vehicle control mice with endometriosis was seen for
195 total moving distance, total moving speed, and peripheral distance, but not for total moving or climbing time,
196 for which no difference between groups was detected. Together, these data indicate that KMO inhibition with
197 KNS898 results in an improvement in well-being evidenced by improved cage exploration behaviour in addition
198 to improved objective histological measures of endometriosis disease burden.
199

200 **DISCUSSION**

201 In this study, we set out to investigate the potential for KMO inhibition as a non-hormonal therapy for
202 endometriosis. First, we confirmed that KMO was expressed in human endometrium by immunohistochemistry,
203 and then showed that KMO was clearly expressed in the epithelial cells in human endometriosis lesions. Next,
204 we demonstrated that the highly specific KMO inhibitor KNS898 was orally bioavailable when given twice
205 daily by gavage in mice, and clearly blocked KMO activity in a dose-dependent manner at a dose of 25mg/kg.
206 We therefore used that dose to test the efficacy of KMO blockade with KNS898 in mice with experimentally-

207 induced endometriosis. One important finding of this project is that KMO blockade resulted in a reduction in
208 endometriosis severity compared to untreated mice with endometriosis, specifically in terms of reducing i) the
209 number of mice that developed endometriosis tissue lesions, and ii) the number of lesions per mouse in those
210 that did develop lesions. The histopathology of the experimental endometriosis lesions was sufficiently similar
211 macroscopically to that seen in the examined human DEGLS, and KMO was evidently highly expressed in the
212 same tissue distribution in model lesions compared to human disease. KMO blockade also decreased lesion
213 KMO expression. Critically, and importantly from a translational perspective, therapeutic blockade of KMO
214 improved visceral hyperalgesia measured by reduced mechanical allodynia and restored normal cage
215 exploration behaviour and mobility in treated mice compared to untreated mice with endometriosis. Together,
216 these data show that KMO is expressed in human and mouse endometriosis tissue lesions and that therapeutic
217 KMO blockade reduces the number of endometriosis lesions and improves holistic metrics of disease behaviour
218 in mice.

219 The model of endometriosis, using inoculation of endometrial tissue of ovariectomized donor mice,
220 reliably induced the pathophysiological symptoms indicative of endometriosis in recipient mice. Test groups
221 inoculated with endometrial tissue (G3-G5) showed significant growth of ectopic endometrial tissue. Groups
222 treated with test article experienced significantly less DEGLS development (significantly fewer DEGLS were
223 noted in treated groups when compared to vehicle treated groups). Disease burden in the treatment group that
224 had treatment starting immediately after inoculation was lower than that of the vehicle-only treated group. It is
225 not clear why mean cystic size and cystic volume in treated animals was not smaller in treated animals. We can
226 only speculate that KMO blockade may potentiate rapid involution of cysts, but this cannot be proven
227 mechanistically here.

228 Mice that received inoculated endometrial tissue showed a measurable and increased visceral
229 hyperalgesic pain response (lower mechanical threshold) as measured by bladder response to von Frey filament
230 testing, and improvement in a surrogate marker of central sensitisation to pain measured by hind paw Von Frey
231 filament testing when compared to control mice. The mechanical threshold of both treatment groups trended
232 higher compared to the vehicle treated group. One interpretation of these data is that KMO inhibition reduced
233 responses to pain caused by the presence of endometriosis, i.e. improving visceral hyperalgesia.

234 Home cage behaviour using HCA indicated a reduced overall activity in endo-inoculated mice when
235 compared to control mice. It should also be noted that test article-treated animals in both groups showed more
236 locomotor behaviour and a seemingly better quality of life within the home cage environment when the home-
237 cage dynamics were monitored. This supports the notion that treated mice exhibit less propensity for behaviours
238 that are, at times, typical of depressive and anxiety-like behaviour in home-cage, group housed conditions.

239 Under the experimental conditions imposed, treatment with KNS898 on two dosing schedules provided
240 a significant reduction in DEGLS formation within the inoculated mice, as well as a seemingly higher pain
241 threshold. This attenuation of pain response was coupled with increased activity levels in the home cage. Taken
242 together, these data suggest a therapeutic effect to alleviation of certain salient symptoms of endometriosis, as
243 well as a reduction in the number and size of DEGLS.

244 Non-pathological endometrium is a site of high KMO expression. Because endometriosis lesions in
245 women are 'endometrial-like' tissue rather than normal endometrium, we tested, and demonstrated expression of
246 KMO in the epithelial layers of endometriosis lesions sampled from women undergoing surgery for

247 endometriosis. First-line medical treatment for endometriosis is the contraceptive pill or other ovarian steroid
248 hormone suppressive drugs. Treatment failures are frequent, side effects are common, all approaches are
249 contraceptive. Many women opt for invasive surgery to remove or ablate the endometriosis lesions.

250 In conclusion, KMO is expressed in human endometriosis tissue lesions and in a mouse model of
251 endometriosis in the epithelial layers of distended endometrial gland-like structures. Oral KNS898 reliably
252 induced a biochemical state of KMO blockade with accumulation of kynurene, diversion to kynurenic acid
253 and ablation of 3-hydroxykynurene production. KMO blockade improved histological and symptomatic
254 behavioural endometriosis disease features with an overall benefit, even when treatment commenced one week
255 after establishment of the disease. KMO blockade is therefore a promising avenue for a new non-hormonal
256 therapeutic modality for endometriosis.

257

258 MATERIALS AND METHODS

259 **Ethical approvals and permissions.** The human tissue samples were obtained from participants who had given
260 fully informed written consent under ethical approval granted by Lothian Research Ethics Committee (LREC
261 11/AL/0376). Human tissue samples were obtained with ethical approval and fully informed consent from
262 individuals attending the Royal Infirmary of Edinburgh as described below. Animal experiments conducted by
263 NAASON Inc were carried out according to the National Institute of Health (NIH) & National Institutes of
264 Health Korea (NIHK) guidelines for the care and use of laboratory animals and approved by Naason Science in
265 accordance with all applicable FELASA, IACUC and AAALAC guidelines. Animal experiments outsourced to
266 Syneos Health were conducted with institutional ethical approval.

267

268 **Human Patients and Samples.** Tissue samples were collected from patients undergoing a diagnostic
269 laparoscopy for suspected endometriosis following Endometriosis Phenome and Biobanking Harmonisation
270 Project (EPHect) guidelines¹³. Patient summary characteristics are presented in **Supplementary Table S1**. Note
271 there was a range of disease stages assigned at time of surgery according to American Fertility Society (AFS)
272 criteria¹⁴. Cycle stage was determined by measuring hormones in blood according to standard protocols and
273 assessment of eutopic endometrial tissue histology when such samples were available¹⁵. Lesions were recovered
274 from 17 patients, of these n=10 were recovered from the peritoneal side wall consistent with classification as
275 superficial peritoneal endometriosis lesions and n=5 from the cysts of ovarian disease (endometrioma) (n=2 on
276 hormones). Eutopic endometrium was from 4 patients n=3 of which had no lesions at time of surgery (noted as
277 stage 0). General histology of samples was assessed using H&E staining.

278

279 **Immunohistochemistry (human endometrium and human endometriosis tissue lesions).** 5 µm sections of
280 formalin-fixed paraffin-embedded tissue blocks were mounted on SuperFrost Plus adhesion slides (Thermo
281 Fisher Scientific). Sections were deparaffined with xylene and rehydrated prior to heat-induced antigen retrieval
282 using Instant Pot: Tris-EDTA pH9¹⁶. Sections were washed with tap water and incubated in phosphate buffered
283 saline (PBS) for 5 minutes. Endogenous peroxidase was blocked with 0.3% hydrogen peroxide in 70% v/v
284 methanol for 30 mins at room temperature then washed in PBS prior to blocking in Normal Goat Serum
285 (NGS)/PBS/bovine serum albumin (BSA)(5%) for 30 mins and streptavidin for 15 mins. Sections were washed

286 twice in PBS and then blocked with biotin for 15 mins and washed in PBS. The primary antibody to KMO
287 (KMO Rabbit polyclonal, Proteintech, Catalog Number:10698-1-AP)¹⁷ was diluted to a final concentration of
288 1:1000 in NGS/PBS/BSA and incubated overnight at 4°C in a humidity chamber. Sections were washed twice
289 with PBS/Tween 0.05% (1ml Tween in 2L PBS) for 5 mins. The secondary detection antibody Goat Anti-Rabbit
290 Biotinylated (Vector Cat number: BA-1000) was diluted in NGS/PBS/BSA (1:500) and incubated for 30 mins,
291 prior to washing twice in PBS/Tween 0.05%, for 5 mins before adding the detection system reagent
292 Streptavidin-HRP (DAKO Cat Number P0397) 1:500 in PBS for 30 min, washed and stained with DAB (DAKO
293 Cat Number K3468) as per manufacturer's directions and incubated for 5 mins before a final wash with tap
294 water. Sections were counterstained with Haematoxylin, dehydrated through graded ethanol and mounted.
295 Sections of stained slides were scanned on a Zeiss Axioscan Z1 slide scanner and exported as TIFF files: images
296 were evaluated for stromal, epithelial and immune cell content.
297

298 **KNS898 preparation for oral administration.** KNS898 powder was weighed and dissolved at the required
299 concentrations in a final vehicle of 2% DMSO, 20% PEG200, 78% 0.15M NaCl by volume. Brief sonication on
300 ice was done to facilitate dissolution.

301
302 **In vivo confirmation of KMO inhibition by KNS898 in mice.** This experiment was outsourced to Syneos
303 Health (Les Templiers, 2400 route des Colles, 06410 Biot, Sophia-Antipolis, France). A formal
304 pharmacokinetic/pharmacodynamic study was not required at this stage. Female C5BL/6J mice aged 12 weeks
305 were purchased from Charles River Laboratories, maintained on standard 12 hour light-dark cycle, given free
306 access to water and standard chow before being randomised to one of three dose levels of KNS898 (n=3 per
307 group, total n=9 mice). Dose levels tested were 0.01mg/kg, 5mg/kg and 25mg/kg. Mice were gavaged with 0.5
308 mL of drug in vehicle twice daily for 7 days before euthanasia and plasma sampling.
309

310 **Plasma samples.** Blood was sampled into Sarstedt Microvette CB K2EDTA 300 µL tubes and centrifuged at
311 5,000 rpm (2380 RCF) for 3 mins. Plasma was aliquoted, frozen on dry ice and transferred to storage at -80°C
312 prior to temperature-controlled shipping.

313
314 **LC-MS/MS analysis of plasma drug levels and kynurenone metabolites.** Plasma samples (100 µL) were
315 diluted at a 1:1 ratio with 4% phosphoric acid and enriched with 50 ng ¹³C₆-kynurenone, ¹³C₆-3-
316 hydroxykynurenone (Sigma Aldrich, custom synthesis) and d5-kynurenic acid (CDN isotopes). 12-point
317 calibration standards (0.1 to 100 ng) were prepared for KNS898, kynurenone (KYN), kynurenic acid (KA), and
318 3-hydroxykynurenone (3HK) and extracted alongside samples using solid phase extraction plates (Waters Oasis
319 HLB, 10 mg sorbent, 30 µm particle size). Extracts were dried down under nitrogen and reconstituted in LC-MS
320 grade water (100 µL). 10 µL was injected onto a column (Ace C18-PFP column; 100 x 2.1 mm internal diameter
321 1.7 µm; HiChrom (VWR, Lutterworth)) using an Acquity I-Class UPLC liquid chromatography system (Waters)
322 linked to a QTRAP 6500+ mass spectrometer (AB Sciex)¹⁰. The flow rate was set at 0.4 mL/min with a column
323 temperature of 40°C. Separation was carried out using a gradient mobile phase system of A – 0.1% aqueous
324 formic acid and B – 0.1% formic acid in methanol, starting at 15% B, rising to 85% B over 6 mins and returning
325 to 15% B by 9 mins. Mass spectrometry settings were for positive mode electrospray (5.5 kV, 700°C) and

326 multiple reaction monitoring m/z 209.0 → 192.2 for KYN, m/z 189.9 → 144.1 for KA m/z 225.0 → 208.0 for
327 3HK and m/z 361.1 → 120.1 for KNS898 and for internal standards were m/z 231.0 → 214.0 for $^{13}\text{C}_6$ -3HK, m/z
328 195.1 → 177.2 for d5-KA and m/z 215.0 → 197.8 for $^{13}\text{C}_6$ -kynurenine. Retention times for KYN, KA, 3HK and
329 KNS898 were 1.8, 3.2, 1.2 and 6.5 mins, respectively and 3.2 mins for d5KA, 1.8 mins for $^{13}\text{C}_6$ -3HK and 1.2
330 mins for $^{13}\text{C}_6$ -KYN. Data were acquired by Analyst 1.7.1 software (AB Sciex) and linear regression analysis
331 was carried out on MultiQuant 3.0.3 software (AB Sciex) where peak integrations and amounts of each
332 kynurenine metabolite and KNS898 were calculated using the peak area ratio of compound/internal standard,
333 with data further handled in Microsoft Excel 2016 as described¹⁸.

334

335 **Experimental mouse model of endometriosis.** This experiment was outsourced to Naason Science Inc.,
336 Osong, Korea (KBIO New Drug Development Center #506, Chungbuk, Korea, 28160) using protocols
337 originally developed by the Saunders team in Edinburgh^{19,20}. Experimental design and groups are shown in
338 **Figure 3a** and **Supplementary Table S2**. There were 5 groups of mice with n=10-15/group):25 mg/kg KNS898
339 was administered twice a day via oral gavage in two of the groups of mice. Group 4 received KNS898 from the
340 time of endometrial tissue inoculation (Day 19; G4); group 5 commenced dosing 1 week after inoculation (Day
341 26; G5). Group 3 received vehicle (2% DMSO, 20% PEG200, 78% 0.15M NaCl) in the same regimen.

342 To perform the mouse model of endometriosis, donor female C57Bl/6 mice aged 6 weeks were acclimatized for
343 2 weeks prior to surgery. Ovariectomy (Day 0) was performed at 8 weeks of age under general anaesthesia with
344 monitoring, with analgesia that extended to the post-operative period with buprenorphine (0.03 ml) (Veterges
345 ic® 3 mg/ml, Ceva Inc., Korea) subcutaneously. To prepare donor tissue that would best replicate menstrual-like
346 tissue in women, ovariectomized (OVX) mice were primed with daily s.c. injections of 100 ng 17 β -estradiol
347 (E2) on days 7, 8 and 9. On days 13 – 19 a silastic progesterone (P4) pellet was implanted subcutaneously.
348 These animals were injected once daily with E2 (5 ng in sesame oil) on days 13, 14 and 15. Decidualization was
349 induced in one uterine horn with an injection of 20 μ l sesame oil 4 hours after the last E2 injection. On day 19
350 (4 days after induction of decidual response), donor mice were killed 4 hours after removal of the P4 pellet.
351 Endometrial tissue was then scraped from the myometrial layer of the decidualized uterine horn, suspended in
352 500 μ l PBS and injected via a XG needle sprayed into the lower abdominal cavity of the recipient mouse under
353 general anaesthesia with monitoring and post-operative analgesia as described. The ratio of donor to recipient
354 mouse was 1:1 (from one donor to one recipient). Recipient mice had intact ovaries to ensure ongoing hormonal
355 stimulation of the injected tissue: group allocations are shown in **Supplementary Table S2**.

356

357 **Mechanical allodynia test by the Von Frey method.** Abdominal and hind paw Von Frey tests were performed
358 in the recipient animals before inoculation (baseline), and 1, 2, and 3 weeks after inoculation. Mechanical
359 threshold was measured using Von Frey filaments. For the hind-paw, 15, 8, 6, 4, 2, 1.4, 0.6, 0.4 g filaments were
360 used, and for the bladder reflex to filament application to the lower abdomen, 60, 26, 10, 8, 6, 4, 2, 1 g filaments
361 were used. The experimenter was blind to the group allocation in order to reduce bias.

362

363 **Cage exploration and behavioural assay using Home Cage Analysis.** Home Cage Analysis (HCA) was
364 performed in the recipient animals before inoculation (baseline), and at the late stage of treatment. Recipient
365 mice were randomly housed using a Monte Carlo randomization. All animals had a micro-chip (BioMark, USA)

366 inserted to the abdomen prior to being placed in the home-cage. This procedure does not cause undue
367 discomfort or hamper, in any way, animal movement. Total moving distance, total moving time, moving speed,
368 isolation/separation distance, isolated time, peripheral time, peripheral distance, in centre zones time, in centre
369 zones distance, climbing time, and body temperature were tracked automatically by an ActualHCA™ Home
370 Cage Analyzer (ActualAnalytics Ltd., Edinburgh, UK) and processed with proprietary machine learning and
371 artificial intelligence algorithms.

372

373 **Endpoint tissue and plasma sampling.** On experimental Day 40, all recipient mice were euthanized, and blood
374 collected via cardiac puncture. Whole blood was collected into heparinised tubes, and plasma was separated by
375 centrifugation (3000 rpm for 15 min) at 4°C. Separated plasma was collected in Eppendorf microtubes, frozen
376 on dry ice and stored at -80°C. Photographs of the abdominal cavity were obtained. Lesions from the abdominal
377 cavity were harvested. DEGLS were dissected from the surrounding abdominal tissue and measured for size and
378 volume. DEGLS volume was measured using the following formula²¹: Volume = long diameter × (short
379 diameter/2)² × π. DEGLS were fixed in a 4% paraformaldehyde solution and prepared for standard H&E. If
380 more than one DEGLS was present in an animal, DEGLS that were not used for H&E were snap-frozen in
381 liquid nitrogen and stored at -80oC.

382

383 **Endometriosis lesion histology.** Endometriosis lesion tissue blocks were sectioned at a uniform thickness of 5
384 μm and were mounted onto a microscope slide. The slide then underwent deparaffination and hydration.
385 Paraffin was removed from the slide using xylene, then hydration through graded ethanol and washing were
386 performed. Slides were then stained with Harris haematoxylin and alcoholic eosin Y and mounted after
387 dehydration and clearing with xylene. The H&E-stained images were visualized using a Slide Scanner
388 (Panoramic scan, 3D HISTECH).

389

390 **Immunohistochemistry (mouse DEGLS).** Immunohistochemistry to detect KMO in mouse DEGLS tissue was
391 performed on a Leica Bond III automated immunostaining robot. 5 μm thick sections obtained from FFPE
392 (formalin Fixed Paraffin Embedded) samples mounted on superfrost plus slides were stained as follows. Heat
393 induced epitope retrieval (HIER) was performed using Epitope Retrieval Solution 1 (Leica, ER1 pH 6.0 citrate
394 based solution) for 20 minutes at 99°C. Tissue sections were then incubated for 10 minutes in hydrogen peroxide
395 to block endogenous hydrogen peroxidase activity followed by 10 minute blocking with normal goat serum. The
396 primary antibody against KMO (Proteintech 10698-1-AP @1:1000 Rabbit)¹⁷ was incubated for 1 hour, then
397 incubated with a goat anti-rabbit peroxidase conjugated secondary antibody for 30 minutes prior to visualisation
398 with diaminobenzoate (DAB) using standard protocols.

399

400 **Digital slide scanning.** Whole sections were scanned using a Zeiss Axioscan Z1 whole slide scanner. The image
401 files (.czi) were batch converted to Tif format for image analysis and quantification, acquisition and batch export
402 used Carl Zeiss Zen v2.5 software.

403

404 **Quantitative image analysis.** Image analysis was carried out using QuPath v0.4.2. Analysis was standardised
405 by using multiple regions of interest from individual slides and collating them into a training image. To improve

406 stain contrast, overlapping DAB and haematoxylin staining was deconvoluted by manual optimisation of the
407 stain vectors. Individual slide epithelia were annotated, with relative DAB optical densities and annotation shape
408 measurements taken. Epithelial thickness and area were calculated to enable correlation with KMO intensity. To
409 generate the heat map of cell KMO expression, a cell detection was carried out on haematoxylin staining using
410 standard parameters, a 5-100um² area range, cell expansion of 1um and a threshold of 0.14.

411

412 **Statistical analysis.** Power calculations were performed using G*Power (v3.1.9.4) software. Input parameters
413 were used: 2-tailed, α -error probability = 0.05, and power (1- β error probability) = 0.80. Continuous variable
414 data were tested for Normality of distribution with a one sample Kolmogorov-Smirnov test. Normally
415 distributed data were analysed by one-way ANOVA with post-hoc Dunnett's T3 for multiple groups. Data
416 comparing treatment group effects at multiple time-points were analysed by two-way ANOVA with multiple
417 comparison testing by Dunnett's method . Data comparing multiple groups with unequal variances were
418 analyzed with Welch's ANOVA. Data not following the Normal distribution were analysed with non-parametric
419 Kruskal-Wallis test. Categorical and proportions data were analysed by Fisher's exact test. Data were visualised
420 with GraphPad Prism.

421

422 **LIST OF SUPPLEMENTARY MATERIALS**

423 Fig. S1 and Fig. S2.

424 Table S1 and Table S2.

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499

500

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510 **AUTHOR CONTRIBUTIONS**

511 Conceptualization: DJM, SPW, PTKS, AH

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513 Investigation: BH, IS, XZ, FC, SGP, JPS, LB, HYL, YGK, KHP, LCP GF, AT, DC, TA

514 Visualization: BH, IS, KHP, LCP, DJM

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519 Writing – review & editing: BH, IS, XZ, FC, NZMH, SGD, JPS, MM, LB, LCP, PJS, AT, SPW, AH, PTKS,

520 DJM.

521

522 **DECLARATION OF INTERESTS**

523 The following authors have interests to declare: S.P.W., D.J.M. are co-founders of Kynos Therapeutics Ltd..

524 D.J.M. is a Board Member of Kynos. The University of Edinburgh controls Patents WO2015/091647,

525 WO2016/097144, WO2016/188827 that relate to inhibitors of KMO inhibitors, and include the compound used

526 in this paper. The remaining authors declare no competing interests.

527

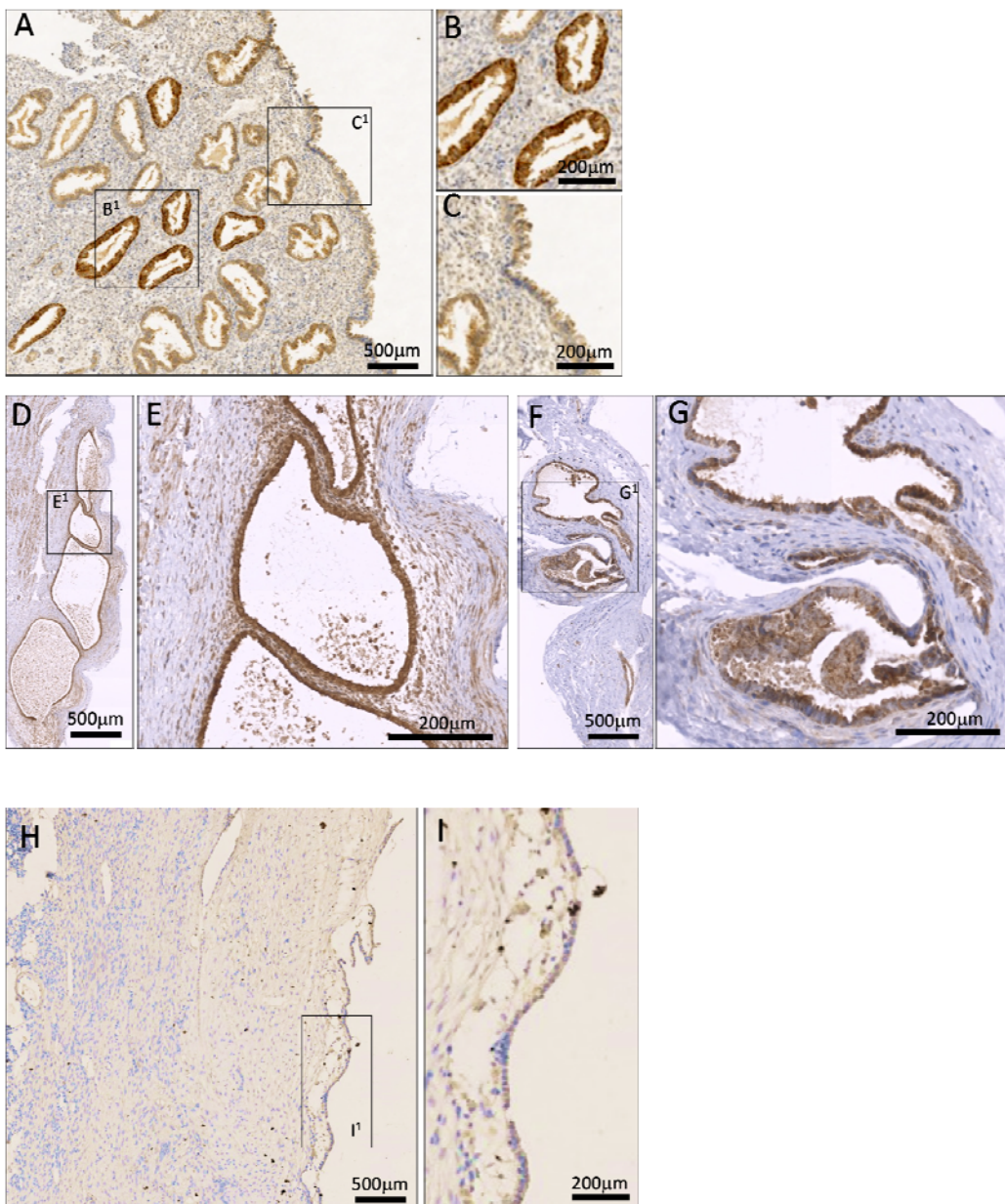
528 **DATA AND MATERIALS AVAILABILITY**

529 All data are available in the main text or the supplementary materials.

530 KNS898 availability is restricted under a Material Transfer Agreement. Please contact the corresponding author
531 in the first instance.

532 **FIGURES**

533 **Fig. 1**

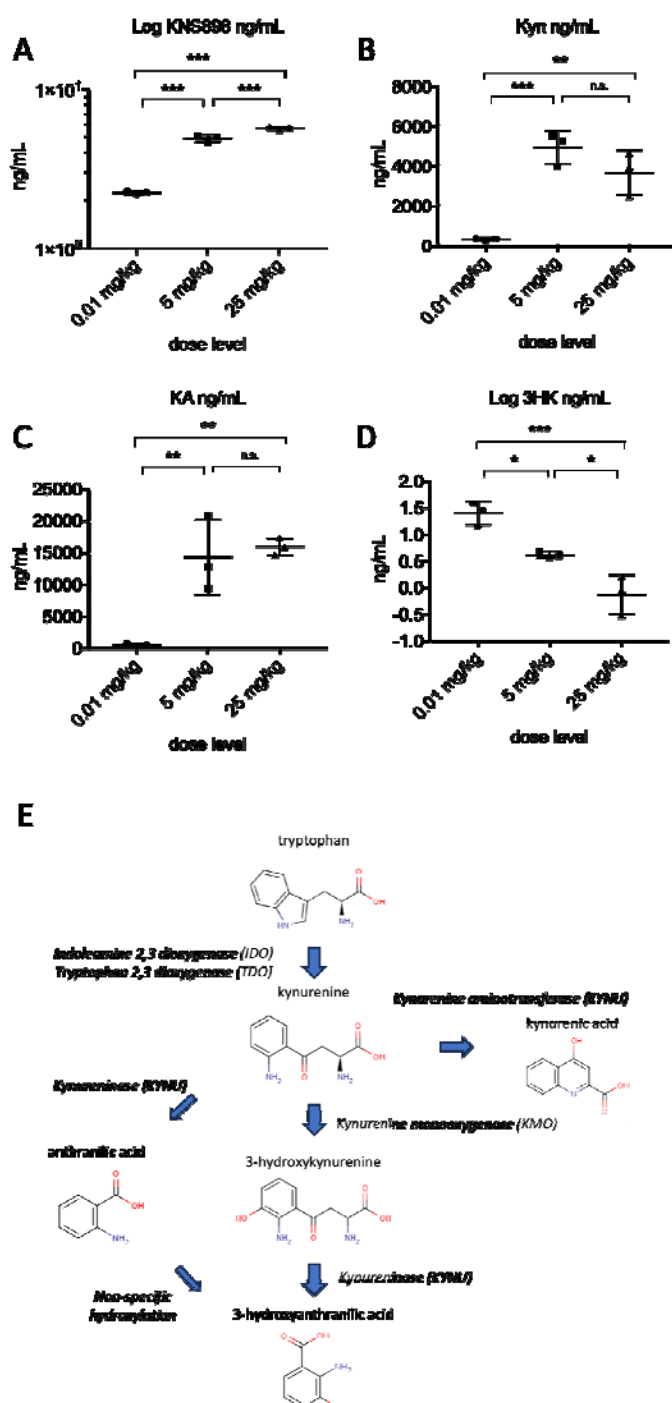


534

535 **Figure 1. Immunohistochemistry of KMO expression in human endometrium and distended**
536 **endometriosis gland-like lesions.** Fixed tissue sections were stained with anti-KMO antibody (1:500 dilution)
537 and visualized with DAB as described in the Methods section. **Panel A.** Normal human endometrium (200 X
538 magnification); **B¹** and **C¹** insets denote areas shown in panels **B** and **C** at higher magnification. KMO
539 expression is demonstrated as dark brown DAB-positive staining and is most intense in the epithelial cells lining
540 the glands. **Panels D through G.** Human peritoneal endometriosis tissue lesions stained with anti-KMO
541 antibody visualized with DAB. D/E PIN3652, Stage II, F/G PIN3306 Stage I. Note intense staining of cells
542 lining glandular structures. **Panel H.** Ovarian-type endometriosis tissue lesion with higher magnification inset

543 (I¹) showing KMO expression present but at lower intensity in the mesothelial tissue surface (this sample does
544 not have epithelial cells in the lesion).

545 **Fig. 2**

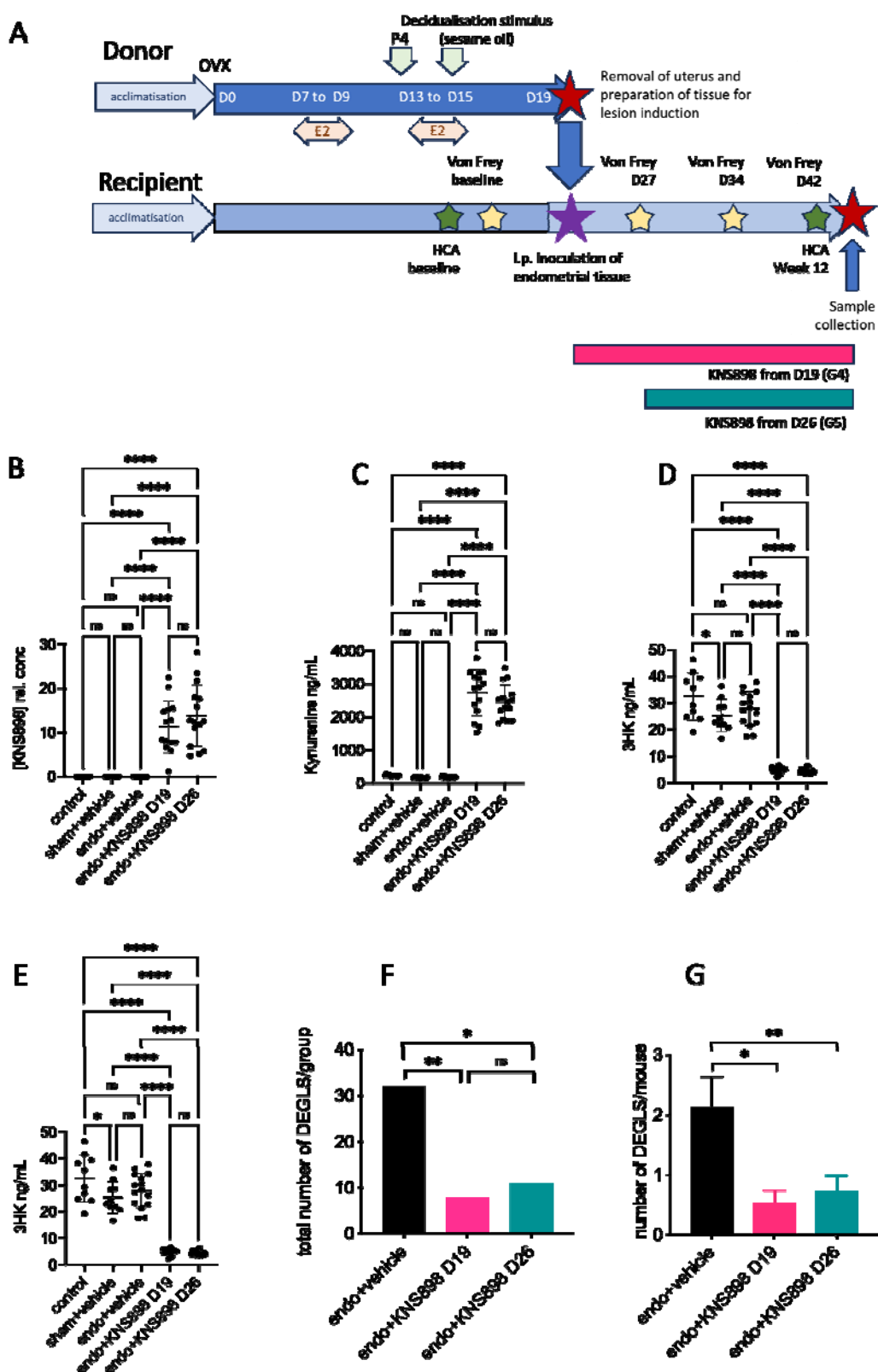


546

547 **Figure 2. KNS898 plasma levels and pharmacodynamic effect of KMO blockade.** Mice (n=3 per group,
548 individual data shown) were given KNS898 twice daily by gavage at the doses shown. After 7 days, blood was
549 sampled at euthanasia and KNS898 levels and kynurenine pathway metabolite levels measured by LC-MS/MS.
550 **A.** KNS898 drug levels. **B.** Kynurene. **C.** Kynurenic acid. **D.** 3-hydroxykynurene (logarithmic scale).
551 Comparison between groups by one way ANOVA with post hoc Tukey's test. *P <0.05, **P<0.01, ***P<0.001,

552 n.s. not statistically significant. **E.** A diagram of the kynurenine pathway showing the key step catalyzed by
553 KMO.

554 **Fig. 3**

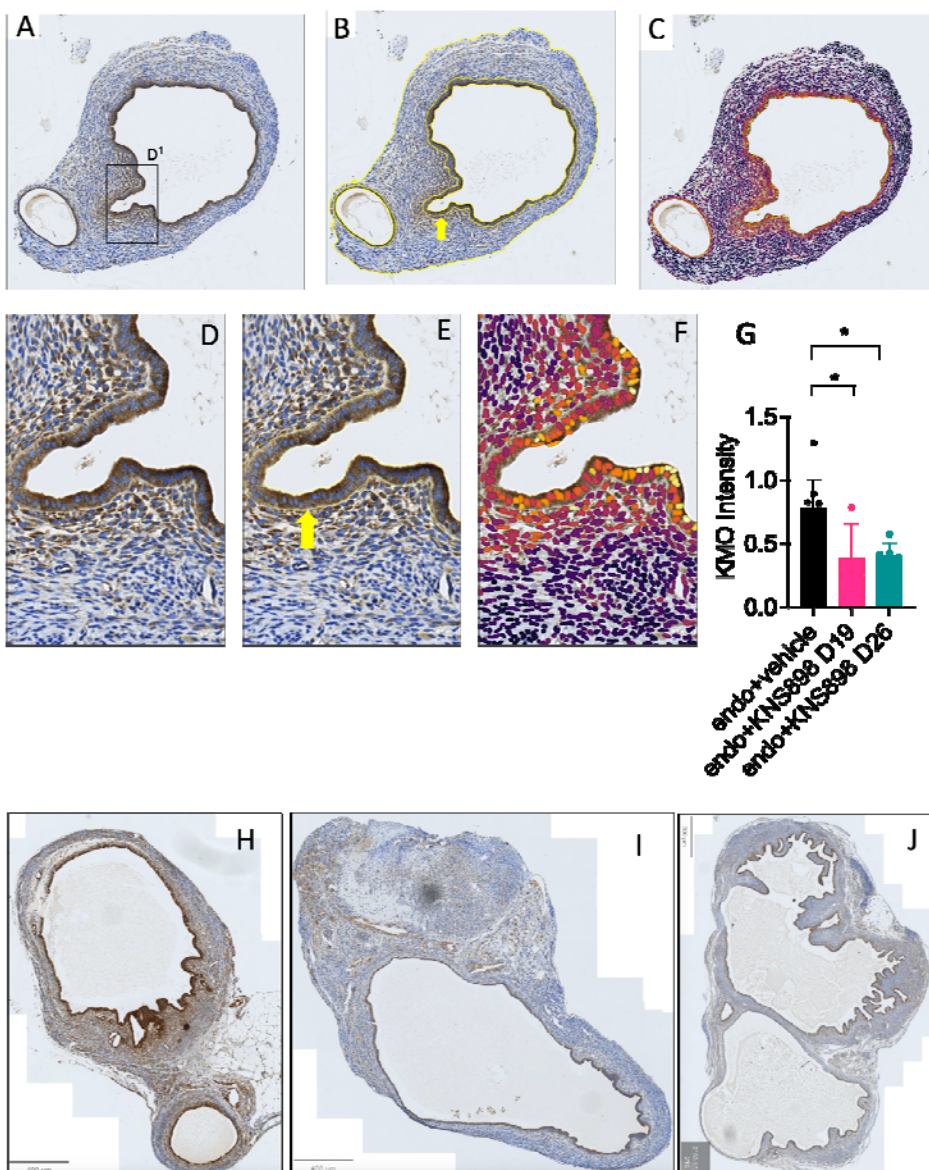


555

556 **Figure 3. Therapeutic effect of KNS898 in an experimental mouse model of endometriosis. A.**
 557 Experimental design. Ovariectomized (OVX) donor mice were hormonally stimulated as shown (E2 estradiol,

558 P4 progesterone). At Day 19, donor mouse endometrial fragments were inoculated into recipient mice in a 1:1
559 ratio. KNS898 treatment 25 mg/kg twice daily by oral gavage was commenced at Day 19 or after a 1 week
560 interval on Day 26 and in both cases continued for 2 weeks. Groups were G1: n=10, control mice; G2: n=10,
561 sham-operated control mice; G3: n=15, endometriosis + vehicle; G4: n=15, endometriosis with KNS898
562 commenced at Day 19; G5: n=15, endometriosis with KNS898 commenced at Day 26. **B.** KNS898 drug levels
563 (relative concentrations). **C.** Kynurenine. **D.** 3-hydroxykynurenine. **E.** Kynurenic acid. Individual data are
564 shown in panels B through E. **Panels F and G.** Enumerated distended endometriosis gland-like structures
565 (DEGLS) in recipient mice by treatment group. **F.** Total number of DEGLS per group. **G.** Total number of
566 DEGLS per animal for all animals in the group; bars show mean with s.e.m. **(G).** Comparison between groups
567 by one way ANOVA with post hoc Tukey's test. *P <0.05, **P<0.01, ***P<0.001, ****P<0.0001, n.s. not
568 statistically significant.

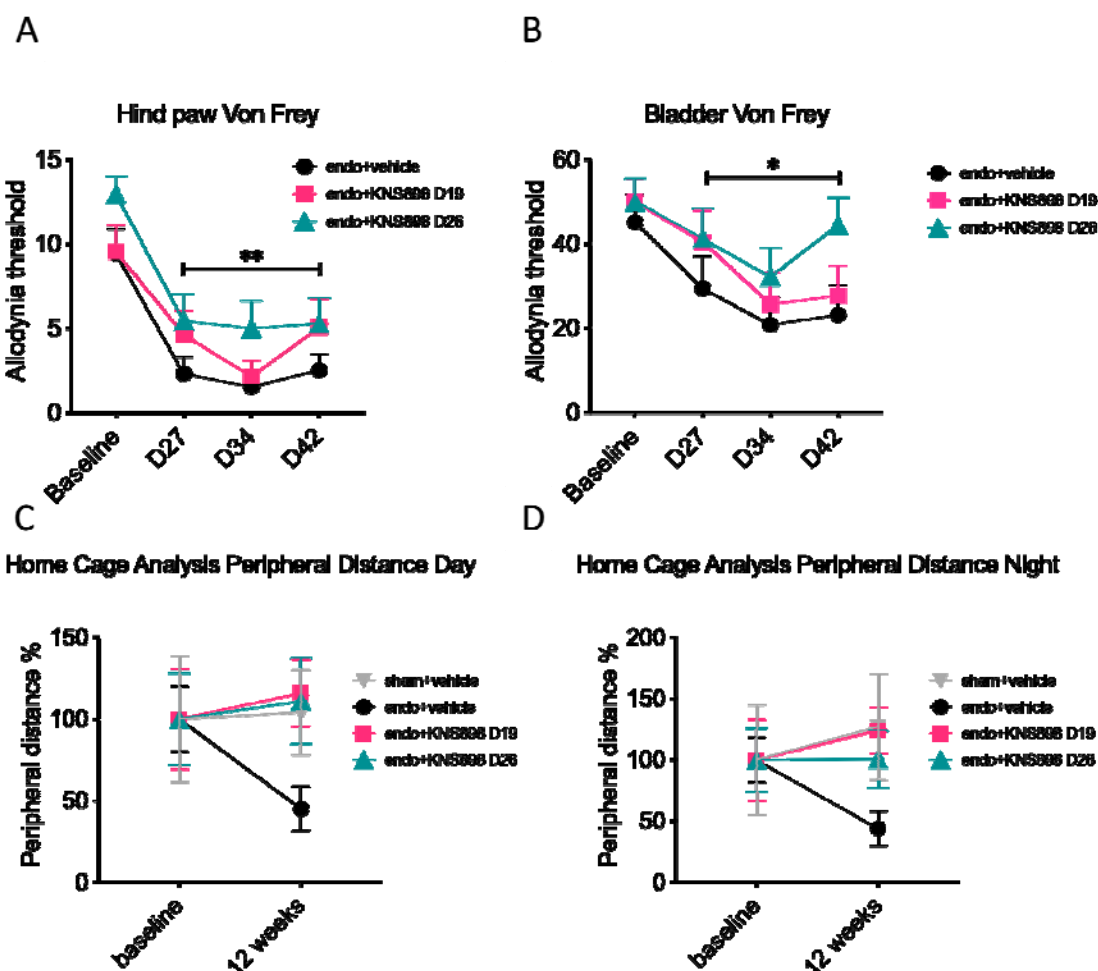
569 **Fig. 4**



570

571 **Figure 4. Quantification of KMO expression in mouse model distended endometriosis gland-like structure**
572 **(DEGLS) lesions.** Sections were visualized at 200 X magnification (A, B, C) with higher magnification insets
573 shown (D, E, F). **Panel A and D.** Fixed tissue sections were stained with anti-KMO antibody (1:500 dilution)
574 and visualized with DAB as described in the Methods section. **B and E.** QuPath was used to identify the
575 epithelial tissue layers (yellow arrows denote the boundary) which were quantified by thickness. **C and F.**
576 KMO expression intensity quantified and heat map expression values are overlaid. **G.** KMO expression
577 staining intensity per unit area of endometriosis DEGLS epithelium, categorized by treatment group (G3
578 endometriosis + vehicle; G4 endometriosis + KNS898 from D19; G5 endometriosis + KNS898 from D26.
579 Individual data points shown. Comparison between groups by one way ANOVA with post hoc Tukey's test. *P
580 <0.05. . **H, I and J.** Representative micrographs from G3 (H), G4 (I) and G5 (J) showing KMO expression in
581 the epithelial cells lining each DEGLS.

582 Fig. 5



583

584 **Figure 5. Effect of KNS898 on mechanical allodynia and illness behaviour in an experimental mouse**
585 **model of endometriosis. A. Hind paw Von Frey filament test showing effect of endometriosis and KNS898**
586 **treatment in groups G3, G4 and G5 B. Bladder Von Frey filament test C. Home Cage Analysis of motility**
587 **showing a daytime motility deficit in mice with endometriosis compared to control mice, and clear restitution of**
588 **normal motility in KNS898 treated groups. D. Nighttime home cage motility analysis showing the benefit of**
589 **KNS898 treatment on normalizing the motility deficit seen in mice with endometriosis. Data are mean with**
590 **s.e.m. For A and B, statistical comparison between groups was by two-way ANOVA to compare Group effect**
591 **and Time effect, with multiple group comparison using Dunnett's T3 test. Asterisks represent treatment group**
592 **effect statistical significance of the Dunnett's T3 test comparing treatment group G5 to G3 vehicle control *P**
593 **<0.05, **P <0.01. For C and D, Welch's ANOVA with multiple group comparison using Dunnett's T3 test was**
594 **used. Although the ANOVA was statistically significant, the post hoc Dunnett's T3 was not, therefore no**
595 **asterisks are marked.**