1	TNF α induced IKK β complex activation influences epithelial, but not
2	stromal cell survival in endometriosis
3	
4	Vida Kocbek ^{1,2} ; Giovanni Grandi ³ ; Fabian Blank ⁴ ; Carlos Wotzkow ⁴ ; Nick A Bersinger ^{1,2} ;
5	Michael D Mueller ^{1,2} , Satoru Kyo ⁵ , Brett D. McKinnon ^{1,2,*} ;
6	
7	
8	¹ Department of Obstetrics and Gynaecology, Inselspital, Berne University Hospital,
9	Effingerstrasse 102, Berne CH-3010, Switzerland ² Department of Clinical Research,
10	University of Berne, Murtenstrasse 35, Berne CH-3010, Switzerland ³ Azienda Ospedaliero-
11	Universitaria Policlinico, University of Modena and Reggio Emilia, Via del Pozzo 71, 41124
12	Modena, Italy ⁴ Live Cell Imaging, Department of Clinical Research, University of Bern,
13	Murtenstrasse 50, Bern CH-3010, Switzerland ⁵ Department of Obstetrics and Gynecology,
14	Shimane University Faculty of Medicine, Shimane, Japan
15	
16	Running title; IKK β in endometriosis
17	
18	
19	*Correspondence address: Brett McKinnon, Department of Obstetrics and Gynaecology,
20	Inselspital, Berne University Hospital, Effingerstrasse 102, Berne CH-3010
21	brett.mckinnon@dkf.unibe.ch
22	

- 24 <u>Abstract;</u> 25
- Study Hypothesis: Can the activity of the IκB kinase (IKKβ) complex in endometriotic cells
 contribute to endometriotic lesion survival?
- 28 Study answer: There is a constitutive activity of the IKKβ catalytic complex in peritoneal

and deeply infiltrating lesions that can influence epithelial, but not stromal cell viability.

30 What is known already: Endometriotic lesions exist in an inflammatory microenvironment

31 with higher local concentrations of cytokines, such as tumor necrosis factor α (TNF α). TNF α

- 32 stimulates the activation of the IKKβ complex, an important nodal point in multiple signalling
- 33 pathways that influence gene transcription, proliferation and apoptosis. However, few data on
- 34 the regulation of IKK β in endometriotic tissue are currently available.
- Study Design, size, duration: A retrospective analysis of endometriotic tissue from
 peritoneal, ovarian and deeply infiltrating lesions from 37 women.
- **Participants/materials, setting, methods:** Basal and activated (phosphorylated) IKKβ concentrations were analysed by Western blotting and immunohistochemistry. The relationship between the expression and activation of these proteins and peritoneal fluid (TNF α) concentrations, measured via ELISA, was examined. A subsequent *in vitro* analysis of TNF α treatment on the activation of IKK β and the effect on epithelial and stromal cell viability by its inhibition with PS1145 was also performed.
- 43 Main results and role of chance; Levels of the phosphorylated IKK β complex in 44 endometriotic lesions had a significant positive correlation with peritoneal fluid TNF α
- 45 concentrations. Phosphorylated IKKβ complex was more prevalent in peritoneal and DIE
- 46 lesions compared to ovarian lesions. IKKβ was present in both epithelial and stromal cells in
- 47 all lesions but active IKKβ was limited to epithelial cells. TNFα stimulated an increased
- 48 expression of phosphorylated IKK β and the inhibition of this kinase with PS1145
- 49 significantly influenced ectopic epithelial cells viability but not eutopic epithelial cells, or
- 50 endometrial stromal cells.
- 51 Limitations, reasons for caution; In vitro analysis on epithelial cells was performed with
- 52 immortalized cell lines and not primary cell cultures and only low sample numbers were

53 available for the study.

- Wider implications of the findings; The regulation of aberrant signalling pathways
 represents a promising yet relatively unexplored area of endometriosis progression. The IKKβ
- 56 complex is activated by inflammation and is critical nodal point of numerous downstream
- 57 kinase-signalling pathways, including NF κ B, mTOR and BAD. This study shows a
- 58 significant relationship between peritoneal fluid TNF α and IKK β activation in epithelial cells
- 59 that will have significant consequences for the continued survival of these cells at ectopic
- 60 locations through the regulation of downstream pathways.

- 61 Large scale data: none
- 62 Study funding/competing interests; The study was funded by the Swiss National Science
- 63 Foundation (Grant Number. 320030_140774). The authors have no conflict of interest to
- 64 declare.
- 65
- 66
- 67 Key Words; endometriosis, kinase, signalling, IKK, TNF, inflammation, transcription factor,
- 68 DIE, peritoneal, endometrioma

69 Introduction

70 Endometriosis is characterized by the growth of endometrial epithelial and stromal cells 71 outside the uterine cavity. It is an extremely prevalent disease occurring in 10-20 % of women 72 of reproductive age and is accompanied by chronic pelvic pain and subfertility. Although the 73 exact pathogenesis is not yet clear Sampson's theory of transplantation is commonly accepted 74 (Sampson, 1928). This theory proposes that viable endometrial epithelial and stromal cells are 75 refluxed back through the Fallopian tube into the peritoneal cavity during menstruation. Once 76 in this ectopic environment these cells avoid immune detection, invade the underlying 77 mesothelial layer and stimulate a chronic inflammatory response. 78 79 Numerous inflammatory cytokines and chemokines (Borrelli et al., 2013; 2014) are increased

80 in the peritoneal fluid of women with endometriosis, which occurs through the coordinated 81 interaction of the refluxed endometrial and peritoneal immune cells. Refluxed endometrial 82 cells produce and secrete chemokines (Hornung et al., 1997) that attract leukocytes and 83 activated peritoneal macrophages (Halme et al., 1983). The activated macrophages produce 84 inflammatory cytokines, which in turn further stimulate cytokine production by the 85 endometrial cells, creating a feed forward regulatory loop (Lebovic et al., 2001) and the 86 chronic inflammatory environment. This inflammatory environment has the potential to both 87 influence symptomology (McKinnon et al., 2015) and disease progression (Bruner-Tran et 88 al., 2013).

89

90 A chronic inflammatory environment can contribute to endometriotic lesion progression 91 through the activation of a series of intracellular kinase signalling pathways (McKinnon et al., 92 2016). The IkB kinase (IKK) complex represents a significant, early nodal point in many of 93 the kinase signalling pathways. In the nuclear factor (NF)kB signalling pathway the IKK 94 complex removes the inhibitory IkB protein from NFkB allowing translocation into the 95 nucleus and gene transcription (Bonizzi and Karin, 2004) subsequently influencing the gene 96 expression of many cytokine and chemokines, immunoreceptors, cell adhesion molecules, 97 stress response genes, and growth factors (Pahl, 1999). IKK β also interacts with the tubular 98 sclerosis (TSC2) protein in the mammalian target of rapamycin (mTOR) pathway influencing 99 cellular proliferation (Lee et al., 2007) and phosphorylates Bcl2-antagonist of cell death 100 (BAD) pathway suppressing apoptosis (Yan et al., 2013).

101

102 The IKK β complex consists of two catalytic subunits, IKK α and IKK β and one regulatory

103 subunit (IKKy) (Hinz and Scheidereit, 2014). The binding of extracellular tumor necrosis

104 factor (TNF)α to its cell membrane receptor TNFR (Haider and Knöfler, 2009) stimulates the

105 phosphorylation of both IKK α and IKK β and activation of the IKK β complex. TNF α

- 106 concentrations are increased in the peritoneal fluid of endometriosis (Harada et al., 1999) and
- 107 are correlated with the severity of the disease (Bedaiwy et al., 2002) and thus TNFa
- 108 stimulated IKKβ activity may have a significant influence on the endometriotic lesions.
- 109
- 110 At present very little information is available on the expression and activity of this important
- 111 upstream nodal kinase in endometriotic cells. In this study we used a combination of clinical
- 112 and *in vitro* experiments to determine the presence and importance of IKKβ in endometriosis.
- 113 We found that peritoneal fluid TNF α concentrations had a significant positive correlation
- 114 with the activated IKK β complex and that this was most likely due to epithelial cell
- 115 expression. Furthermore IKKβ activity was important in regulating ectopic epithelial cell, but
- 116 not stromal cell survival. These results therefore suggest a significant role of IKK β in
- 117 endometriotic epithelial cells that deserves further attention.

118 Methods;

119 Patient samples

120 Prior to surgery the relevant institutional review board granted ethical approval and informed 121 consent was obtained from all patients. During surgery performed for suspected 122 endometriosis samples of endometrium, peritoneal fluid and endometriotic lesions were 123 collected. Endometrial biopsies were collected via soft curette (Pipelle de Cornier, 124 Laboratoire CCD, France) and stored in RNAlater (Thermo Fischer Scientific, USA) at -80 125 °C as described previously (Santi et al., 2011). Peritoneal fluid was collected during the 126 laparoscopic procedure from the cul-de-sac and centrifuged to remove blood cells, aggregates 127 and debris. Samples were excluded if blood remained in the samples. The pelvic cavity was 128 examined and any endometriotic lesions were removed and the patient staged (no 129 endometriosis, or stage I-IV) according to the revised American Fertility Society staging 130 system (rAFS) ("Revised American Society for Reproductive Medicine classification of 131 endometriosis," 1997). The lesions were recorded as either peritoneal, ovarian or deeply 132 infiltrating endometriosis (DIE). All surgeries were performed during the proliferative phase 133 of the menstrual cycle and endometriosis was confirmed by histological analysis.

134

135 Endometrial biopsies were collected from both women with (n = 8) and without 136 endometriosis (n = 7) and used for the isolation of primary cells via collagenase digestion and 137 size exclusion, as described previously (McKinnon et al., 2012). Strong yields were obtained 138 for the primary stromal cells (ESC) from all women except one without endometriosis, 139 however only limited amounts of epithelial cells could be successfully isolated. As endometrial epithelial cells are terminally differentiated and do not propagate immortalized 140 141 epithelial cells were acquired. Matching peritoneal fluid of sufficient quality was not always 142 available for cytokine measurement therefore if peritoneal fluid was available the ectopic 143 lesions were immediately frozen and stored for stored fresh frozen for subsequent Western 144 blot analysis. If peritoneal fluid was not available, they were formalin fixed and paraffin 145 embedded for immunohistochemistry analysis.

146

147 Cytokine measurement in peritoneal fluid

148 TNFα was measured by an enzyme-linked immunoabsorbent assay (ELISA) kit (R&D
149 Systems, Abingdon, England) using a high-sensitivity NADH cascade amplified alkaline
150 phosphatase with antigen-antibody incubations at 28 °C in a dry incubator and at a dilution of
151 1:2 in the diluent provided. Peritoneal fluid progesterone concentrations were also measured
152 via a radioimmunoassay (Coata-count, DPC; Buhlmann Laboratories, Allschwil, Switzerland)
153 to confirm the patient cycle phase (McKinnon *et al.*, 2014).

155 Protein isolation and Quantification in ectopic lesions

Approximately 30mg of fresh frozen ectopic tissue was used to prepare whole cell extracts via homogenization with the FastPrep 120 tissue homogenizer (30 seconds at 4.0 m/s) in radioimmunoprecipitation assay buffer (RIPA; 50mM Tris-Cl, pH 7.4, 150mM NaCl, 1mM EDTA, 1% v/v triton X-100, 1 % w/v sodium deoxycholate, 0.1% w/v sodium dodecyl sulfate and 1% v/v protease and phosphatase inhibitor cocktail (Cell Signalling Technology, Danvers, Massachusetts)). Final protein concentrations were determined by the bicinchoninic

acid assay (QuantiPro BCA; Sigma).

163

164 Proteins separation was performed by heating 20µg of total protein into LDS buffer 165 (Invitrogen) to 70 °C for 10 minutes and running it on a 4 -15 % Novex NuPAGE Bis/Tris gel 166 (Invitrogen). Proteins were transferred to a 0.45µm nitrocellulose membrane in 4-167 morpholinepropanesulfonic acid buffer (MOPS; Invitrogen) pH 7.7 for 1 hour. Non-specific 168 binding was blocked by incubation with 5% bovine serum albumin (BSA) in phosphate 169 buffered saline (PBS) containing 0.1% Tween 20 (PBST). Membranes were probed with 170 rabbit anti-IKKß antibody (1:1000) (Cell Signalling Technology), and with rabbit anti-171 pIKKa/β(176/180)(1:1000) antibody (Cell Signaling Technology), or mouse anti-actinβ 172 antibody (Abcam, Cambridge, UK) 1:5000. Secondary anti-rabbit (GE Healthcare, Opfikon, 173 Switzerland) and anti-mouse (Sigma) antibodies conjugated to horseradish perioxidase were 174 diluted 1:50 000 and 1:200 000 respectively. Immuno-reactivity was determined with the 175 SuperSignal West Femto kit (Pierce; Thermo Scientific) using the Chemi-Doc XRS+ system 176 (Bio-Rad Laboratories, AG, Cressier, Swituzerland). Band densitometry was quantified with 177 the Quantity One software and in each Western blot a calibrator sample with strong IKKβ, 178 and pIKK α/β expression was included to normalize concentrations across gels and determine 179 protein concentrations relative to the calibrator sample. The relative expression of each 180 sample was expressed as a percentage of the calibrator. Actin β was used as a loading control.

181

182 Immunohistochemistry

183 Immunohisotchemistry was performed using serial sections of 4µm mounted onto glass slides 184 (Superfrost, Braunschweig, Germany), dewaxed in xylene and rehydrated through a series of 185 decreasing ethanol concentrations. Epitope retrieval was performed with 10mM citrate buffer, 186 pH 5.5 for 5 minutes in a 450W microwave. Endogenous perioxidase activity was blocked 187 with 3% hydrogen perioxide (H2O2) and a blocking step performed with 3% BSA for 30 188 minutes in Tris buffered saline (TBS; Tris 100mM, NaCl 0.15 M; pH 7.4). Rabbit anti-IKKß 189 antibody (1:100) and rabbit anti-pIKK α/β (176/180) antibody (1:100) were diluted in 3% 190 BSA in TBS and incubated at 4 °C overnight in a humidified chamber. Slides were washed 191 with TBS and 0.1% Tween 20 (TBST) prior to incubation with an affinity purified, biotin

- 192 conjugated goat anti-rabbit antibody (Dako, Glastrup, Denmark) for 90 minutes at room 193 temperature. After a final wash slides were incubated with an avidin-biotin HRP complex 194 (Vectastain, ABC Kit, Vector Laboratories, Burlingham, CA, USA) for 45 minutes. The 195 antigen-antibody complex was detected by incubation with 3,3' diaminobenzidine substrate 196 and slides were counterstained with hematoxylin and mounted in Aquatex (Merck). The 197 primary antibodies were excluded for the negative controls. Images were photographed with a 198 Nikon Eclipse E800 microscope (Nikon, Japan). Semi-quantitative analysis of antibody 199 staining in the epithelial and stromal cells of the endometriotic lesions was determined by the 200 allocation of scores between 0-3; 0 (negative), 1 (weak), 2 (moderate) and 3 (strong), based 201 on the intensity of staining. The percentage of cells with positive immuno-reactivity was also 202 determined and allocated a score between 0-6 as described 0% = 0, 1-10% = 1, 11-30% = 2, 203 31-50% = 3, 51-70% = 4, 71-90% = 5; > 91% = 6 in each cell type of the endometriotic 204 lesions. For the final immuno-reactive score the staining intensity and percentage of positive 205 cells was multiplied, as described previously (Samartzis et al., 2012).
- 206

207 Cell culture and TNFα treatment

208 Isolated primary endometrial stromal cells (ESC) were maintained in Iscoves's modified 209 Eagle medium (IMEM) (Invitrogen) supplemented with 10% fetal calf serum (FCS) 210 (Invitrogen) and 1% antibiotic/antimycotic (Invitrogen). The immortalized epithelial cell 211 lines, EM E6/E7 and EM'osis, were provided by Professor Kyo, Kanazawa, Japan and were 212 isolated from eutopic endometrium (Kyo et al., 2003) and an ectopic endometrioma (Bono et 213 al., 2012) respectively. These cells were maintained in Dulbecco's modified Eagles medium 214 (DMEM) (Invitrogen) with 10% FCS and 1% antibiotic/antimycotic. The 12Z cells were 215 provided by Professor Starzinski-Powitz, Goethe University and were originally isolated from 216 a peritoneal endometriotic lesion (Zeitvogel et al., 2001). These cells were also maintained in 217 complete DMEM media.

218 To determine the influence of TNF α on pIKK α/β activity in all cell types the cells were 219 seeded into 6 well plates at approximately 3 x10⁵ cells/ well. After reaching approximately 220 80% confluence the media was changed to 0.5% FCS for overnight incubation prior to 221 treatment. Cells were treated either with control media (0.5% FCS in normal media) or 222 control media plus 10ng/ml and 100ng/ml recombinant human TNF α (R&D systems, United 223 Kingdom) for 6 hours. At the end of the treatment period the cells were rinsed and collected 224 in RIPA buffer.

225

226 MTS assay and treatment with PS1145

227 PS1145 is a small molecular weight compound that is a specific inhibitor of IKKβ activity

228 (Lam *et al.*, 2005). Inhibition of IKKβ activity with PS1145 was performed in 96 well plates

229 seeded at a density of $6x10^{3}$ / well. Twenty-four hours prior to treatment the cells were 230 changed into serum free media and treatment media prepared by diluting PS1145 into either 231 serum free media at a final concentration of 1µM. Subsequent concentrations were prepared 232 by a 1:3 serial dilution (333.33nM, 111.11nM, 37.04nM, 12.3nM & 4.12nM). Cell viability 233 was measured after 72 hours by the CellTiter 96 Aqueous One Solution Cell Proliferation 234 Assay (Promega). Triplicate wells were used for each cell type and experiment. For the 235 immortalized epithelial cell lines the experiment was repeated three times and for the primary 236 ESC an experiment on each of the eight endometriosis and seven non-endometriosis 237 preparations was performed separately. A control (without PS1145) was included for each 238 experiment and designated as 100% viability and subsequent values expressed as a percent of 239 control.

240

241 Statistical Analysis;

242 All statistical analyses were performed with Graphpad Prism version 6.0. The correlation 243 between the peritoneal fluid cytokines and IKK\beta and pIKKa/ß expression was performed 244 using the non-parametric Spearman's rank correlation coefficient. Two groups comparisons 245 were performed with a non-parametric Mann-Whitney U test and the comparison of three or 246 more groups with the non-parametric Kruskal-Wallis One-way analysis (ANOVA) and 247 Dunn's multiple comparison Post hoc test. The interaction between two variables was 248 determined via a two-way ANOVA with a post-hoc Tukey's multiple comparison test to 249 determine the difference between individual groups or conditions.

251 Results;

252 Patient data and characteristics

In total, endometriotic tissue was removed from 37 endometriosis patients and endometrial biopsies from eight women without endometriosis. Of these 37 women we collected accompanying peritoneal fluid from 21 in order to compare peritoneal fluid TNFα and ectopic IKKβ. Of the 21 samples 14 were collected from women without any hormonal treatment, four were using combined oral contraceptives (COC) and three were using GnRH analogues. No significant variation in TNFα, IKKβ or pIKKα/β based on hormonal use (**Table 1**) was identified. Five of the lesions were peritoneal, eleven ovarian and five DIE.

260

261 The remaining 16 samples without accompanying peritoneal fluid were kept for 262 immunohistochemistry. Of these, six women had no history of hormonal treatment, five were 263 using COC and five were using GnRH analogues. Three lesions were peritoneal, seven 264 ovarian and six DIE. No significant variation was observed between either pIKK α/β , or IKK β

expression in both the epithelial and stromal cells based on hormonal treatment (Table II).
266

- 267 *IKKβ* expression and activation in endometriotic tissue and its relationship to peritoneal
 268 *fluid TNFα concentrations;*
- 269 Comparison of peritoneal fluid TNF α and endometriotic lesion IKK β and pIKK α/β showed a 270 significant positive correlation between TNFa and pIKKa/ β (r = 0.6268, n = 21, p = 0.0024) 271 (Figure 1A), but not IKK β (r = 0.4216, n = 21, p 0.0570) (Figure 1B), as determined by semi 272 quantitative Western blot (Figure 1C). A significant variation in pIKK α/β concentrations 273 between lesions from different locations (p < 0.05) was observed with a *post-hoc* analysis 274 confirming a significantly lower expression in ovarian lesions (50 \pm 8.7, n =11) compared to 275 the peritoneal lesions $(99 \pm 9.7, n = 5)$ (p = 0.041) (Figure 1D). No significant difference was 276 observed with the DIE lesions (91 \pm 21.6, n = 5). In contrast, IKK β expression showed no 277 variation between lesion types (p = 0.4905) with similar expression in the peritoneal (138 \pm 278 29.3, n = 5), ovarian (128 ± 26.7, n = 11) and DIE lesions (80 ± 26.9, n = 5) (Figure 1E). The 279 active to inactive ratio of IKK β (pIKK α/β :IKK β) varied significantly between lesions (p = 280 0.0123) with both peritoneal $(1.0 \pm 0.39, n = 5)$ and DIE $(1.7 \pm 0.54, n = 5)$ higher than 281 ovarian lesions $(0.5 \pm 0.09, n = 11)$ with a *post-hoc* analysis showing a significant difference 282 between DIE and ovarian lesions (p = 0.0168).
- 283

284 Cell specific activation of IKK in endometriotic lesions

285 Immunohistochemistry identified a low but uniform expression of IKKβ in peritoneal (Figure

286 2A), ovarian (Figure 2B) and DIE lesions (Figure 2C) in both stromal (red arrows) and

287 epithelial cells (*black arrows*). In contrast, pIKK α/β in peritoneal (Figure 2D), ovarian

Brett McKinnon 28.7.2016 22:48
Kommentar [1]: The value in the table
has been corrected

- 288 (Figure 2E) and DIE (Figure 2F) lesions was predominantly epithelial (black arrows), with
- 289 significantly less stromal cells expression (red arrows). Negative controls showed no
- 290 expression in peritoneal (Figure 2G), ovarian (Figure 2H), or DIE (Figure 2I) lesions. No
- 291 statistically significant difference was observed in IKKβ expression between either lesion
- location (p = 0.2420) or cell type (p = 0.1972) (Figure 2J), although this could be due to a
- 293 lack of power. pIKK α/β expression was significantly different in cell types (p = 0.0198), but
- 294 no statistically significant difference could be observed between lesion type (p = 0.3402)
- 295 (Figure 2K) possibly again due to lack of power.
- 296

297 *IKKβ* activity after TNFa treatment298

299 Western blot analysis of pIKK α/β after TNF α treatment of epithelial cell cultures confirmed a 300 low but positive expression in all cell lines examined (Figure 3A). Semi-quantitation of band 301 densitometry indicated that the 12Z cells showed the strongest basal expression (no $TNF\alpha$), 302 but this did not vary after TNF α treatment (p = 0.2320). The EM'E6/E7 cells showed lower 303 basal expression and also no significant variation after TNF α treatment (p = 0.4475). In 304 contrast, TNF α treatment of EM'osis cells significantly increased pIKK α/β above the no 305 treatment control (46 ± 5.0 , n = 3) at concentrations of both 10ng/ml (141 ± 34.2 , n = 3, p = 306 0.0173) and 100ng/ml (197 \pm 23.3, n = 3, p = 0.0085) (Figure 3B). Similar Western blots 307 were performed on protein lysate isolated from stromal cells however no protein expression 308 could be observed in these preparations.

309

310Cell specific influence of IKKβ inhibition on viability311

312 In the epithelial cell cultures there was a significant influence of PS1145 on cell viability 313 based on cell type ($p \le 0.0001$) (Figure 4A). No significant effect of PS1145 was observed on 314 the EM E6/E7 at any concentration. For EM'osis cells there was a significant increase in cell 315 viability at the lowest concentrations (PS1145 2.43nM; 123 ± 0.2 , n = 3, p < 0.001) that was 316 gradually diminished as concentrations increased (PS1145 1 μ m; 106 ± 6.3 p > 0.05), whereas 317 12Z cell viability was significantly decreased at the lowest concentrations (PS1145 2.43nM; 318 70 ± 4.6 , n = 3, p < 0.0001) and remained significantly reduced through to the highest 319 concentration (PS1145 1µm; 73 \pm 2.6, n = 3, P < 0.0001). In contrast primary ESC isolated 320 from women with and without endometriosis showed no significant variation based on either 321 PS1145 (p = 0.8868) or cell type (p = 0.3516) (Figure 4B).

- 322
- 323
- 324

nris Ford 27.7.2016 11:38

Kommentar [2]: Please be wary of interpreting this to mean that the values are similar.

Brett McKinnon 28.7.2016 23:00 **Kommentar [3]:** I have included the phrase 'statistically significant'

325 Discussion

326

327 In this study we examined the expression of the IKK β protein kinase complex and the 328 activation of its catalytic subunits pIKK α/β in both endometriotic lesions, as well as its 329 influence on cell survival in *in vitro* models. The results show a significant relationship 330 between the phosphorylation of the IKK β complex and peritoneal fluid TNF α . Subsequent 331 immunohistochemistry staining showed that although no statistically significant difference in 332 IKK β expression was observed across all cells and lesion types the phosphorylated IKK β 333 complex was predominantly epithelial. This data was supported by the in vitro studies that 334 confirmed pIKK α/β expression in epithelial cell culture models, but not primary stromal cells 335 and that inhibition of IKKB activity significantly influenced endometriotic epithelial cell 336 viability, but not eutopic epithelial cell viability, nor the viability of endometrial stromal cells 337 from women with and without endometriosis. These results therefore suggest that a 338 dysregulation of the IKK β kinase occurs in ectopic epithelial cells that may be related to the 339 inflammatory microenvironment. Given the role of IKKB in transmitting extracellular signals 340 into cell survival via kinase signalling pathways it may represent a significant molecule in 341 endometriosis pathogenesis.

342

343 At present there is very little known about the role of IKK β in endometriosis. The results 344 from our clinical samples suggest that the constitutive IKKß activity is significantly different 345 between peritoneal and DIE lesions compared to ovarian lesions. This difference of 346 expression was supported by our *in vitro* results that showed TNF α stimulated an increase in 347 pIKK α/β expression and that inhibition of IKK β activity increased the Em'osis cell viability. 348 In contrast in the peritoneal derived epithelia cells TNF α had a limited influence of pIKK α/β 349 expression and inhibition of IKKB activity decreased cell viability. Unfortunately a DIE 350 derived cell line was not available.

351

352 Endometriosis is a significantly heterogeneous condition, although whether these lesions have 353 different pathologies (Nisolle and Donnez, 1997), or represent a continuum of the same 354 disease (Somigliana et al., 2004) is still debated. These data suggest a varied cellular response 355 to inflammation may occur in different lesions. It has previously been shown that rectovaginal 356 septum lesions have a distinctly inflammatory phenotype (Bertschi et al., 2013) and that 357 concentrations of inflammatory mediators are stronger in the peritoneal fluid of DIE 358 compared to lesions from other locations (Santulli et al., 2012). As IKKβ can be associated 359 with TNF α both in ours and other studies (Lee *et al.*, 2007) it is possible that the increased 360 production of inflammation associated with DIE lesions is related to the higher IKKβ activity.

is Ford 27.7.2016 11:50

Kommentar [4]: Please consider if this conclusion is justified. Some groups are small so statistical power is limited.

Brett McKinnon 28.7.2016 23:00 Kommentar [5]: I have changed the wording to temper the conclusion 361 The identification of a TNF α influenced activation of the IKK β in endometriotic tissue is 362 significant because of the multiple downstream pathways it regulates (Figure 5) and the 363 influence this can have on gene transcription, protein translation and both cellular 364 proliferation and apoptosis. Activation of IKKβ stimulates NFκB gene transcription and a 365 constitutive activation of NFkB has been observed in peritoneal endometriosis (González-366 Ramos et al., 2007). Multiple factors present in the peritoneal fluid of women with 367 endometriosis including cytokines and iron overload (Alvarado-Díaz et al., 2015) may lead to 368 this constitutive activation. Furthermore an increased NFkB activity has been linked to 369 recurrence of ovarian endometrioma (Shen et al., 2008). Neither of these studies however 370 examined IKKB expression directly. In the immortalized epithelial 12Z cells IKKB inhibition 371 attenuated inflammatory cytokine secretion (Grund et al., 2008) and in ectopic endometrial 372 stromal cells miR200a suppresses IKK β (Dai *et al.*, 2012), raising the possibility of 373 suppressed IKKB activity in stromal cells occurs via an epigenetic regulation. In contrast to its 374 role in inflammation via the NF κ B pathway, IKK β can also regulate cellular proliferation and 375 apoptosis through the mTOR and BAD pathways (Dunlop and Tee, 2014) (Yan et al., 2013). 376 A dysregulation of mTOR has previously been implicated in endometriosis pathogenesis of 377 DIE lesions leading to increased proliferation (Leconte et al., 2011), as has a role for mTOR 378 mediated autophagy (Choi et al., 2014) and BAD activation in ovarian endometriomas 379 (Stickles *et al.*, 2015). Together this suggests that TNF α has the potential to modulate all of 380 these activities via IKKß activation.

381

382 We found that the constitutive activation and influence on cell survival was largely restricted 383 to epithelial cells. Endometriotic lesions are a combination of epithelial and stromal cells and 384 an interdependency between the cells types is required for endometriotic lesions to continue 385 proliferating as tissue integrity of refluxed endometrial matter is essential to endometrial 386 tissue implantation (Nap et al., 2003). We have also previously shown that the stromal cells 387 produce significantly more inflammatory cytokines than epithelial cells in response to 388 stimulation (Bersinger et al., 2008). It could therefore be postulated that a paracrine 389 regulation occurs in the lesions through the stromal cells production of cytokines stimulating 390 a constitutive activation of the IKK^β complex in epithelial cells, which ultimately contributes 391 to cell survival. More research however is required to explore this hypothesis.

392

393 Furthermore, the activity of IKK β in other cell types other than endometriotic cells was not 394 directly addressed in this study, but may give further insight into this mechanism in normal 395 tissue. For endometriosis, however whether this mechanism also happens in healthy eutopic 396 epithelial cells may be of less consequence. This is because epithelial cells will only be 397 present in the peritoneal cavity when endometriosis is present, and when endometriosis is 398 present there is a constant inflammatory environment. We believe it is this confluence of 399 ectopic epithelial cells and constant inflammation that makes the contribution of $TNF\alpha$ 400 stimulated IKK β activity significant. It may be such that this is a characteristic not inherent in 401 the endometrium, but rather acquired during the life of the lesion and contribute more to 402 progression through a constant stimulation of the inflammatory cascade. Further study on 403 whether there is a significant difference between the activation of IKK β in the eutopic 404 endometrium of women with and without endometriosis would be an interesting follow-up.

406 Whether other cell types also show a constitutive activity of IKK β in the presence of 407 inflammation would also be interesting. In this study the images in Figure 2 indicate cells 408 proximal to the endometriotic lesion are largely negative for pIKKa/ß expression, providing 409 circumstantial evidence for the preferential activation of pIKK α/β in endometriotic epithelial 410 cells. Previous studies suggest that cells proximal to the lesion may have different 411 characteristics to cells distal to the lesions (Young et al., 2014) and these cells thus may also 412 be interesting to study, however we were unable to collect this tissue due to our current 413 ethical approvals. Future studies on this topic may however be warranted.

414

405

415 Limitations of this study were the inclusion of women with hormonal treatment. Previous 416 research however has suggested there was no significant influence on NFkB activation by 417 oral contraceptives (González-Ramos et al., 2007). We also observed no statistically 418 significant difference for IKKB in this study, although the power of this analysis was limited 419 by the small sample size. It is possible that this may introduce a variability in peritoneal fluid 420 cytokine concentrations in endometriotic women, as GnRHa analogues have been shown to 421 have an influence on the inflammatory environment (Nirgianakis et al., 2013), however we 422 postulate that a reduction in inflammatory cytokines by hormonal treatment would also be 423 reflected by a subsequent reduction in IKKß activation, maintaining any correlation between 424 the extra and intracellular environment. A further limitation of this study is the small sample 425 size. An expansion of the sample number would provide more definitive information on both 426 the influence of TNF α , as well as hormonal treatments, on IKK β activity. It is difficult to 427 draw direct conclusions on the contribution of IKK β to cell survival in different lesions types 428 as immortalized cell lines were used, however given the similarity observed in the in vitro 429 results to the clinical samples we can be confident that IKK β has a role in epithelial 430 endometriotic cells.

431

432 In conclusion we have observed a significant relationship between TNF α and the activation of 433 IKK β complex in the endometriotic microenvironment and that this activation occurs 434 predominantly in the epithelial cells of peritoneal and DIE lesions. Furthermore IKK β Chris Ford 27.7.2016 12:03

Kommentar [6]: Given the number of observations and the variance, please consider whether hormonal; effects can be excluded with such confidence.

- 435 inhibition in vitro significantly influenced epithelial cell, but not stromal cell behaviour. The
- 436 regulation of kinase signalling pathways is a significant, but under explored area of
- 437 endometriosis pathogenesis and progression and may represent potential non-hormonal
- 438 treatment targets for endometriosis (McKinnon et al., 2016). Given the ability of the
- 439 extracellular inflammatory environment to influence ΙΚΚβ activity and its subsequent affect
- 440 on downstream pathways this kinase may be significant interest in endometriosis.

441

- 443 Acknowledgements; The authors would like to acknowledge the skilful work of Anne
- 444 Vaucher in the running of ELISA measurements. The assistance of the live cell imaging core
- 445 facility of the Department of Clinical Research and funding provided by the Swiss National
- 446 Science Foundation.
- 447
- 448 Authors Roles; VK performed experiments, cell culture and assisted with sample collection.
- 449 CW performed immunohistochemistry and analysis. GG assisted with cell culture. NAB
- 450 assisted with sample collection and intellectual development of the project. MDM assisted
- 451 with sample collection and intellectual development of the project. **BDM** conceived project,
- 452 performed experiments and prepared the manuscript.
- 453
- 454 **Funding;** Support for this project was provided by the Swiss National Science Foundation
- 455 (Grant No. 320030_140774).
- 456
- 457 Conflict of Interest:
- 458 None
- 459

460 461	References; Alvarado-Díaz CP, Núñez MT, Devoto L, González-Ramos R. Iron overload-modulated
462	nuclear factor kappa-B activation in human endometrial stromal cells as a mechanism
463	postulated in endometriosis pathogenesis. Fertil Steril 2015;103:439-447.
464	Bedaiwy MA, Falcone T, Sharma RK, Goldberg JM, Attaran M, Nelson DR, Agarwal A.
465	Prediction of endometriosis with serum and peritoneal fluid markers: a prospective
466	controlled trial. Hum Reprod 2002;17:426-431.
467	Bersinger NA, Frischknecht F, Taylor RN, Mueller MD. Basal and cytokine-stimulated
468	production of epithelial neutrophil activating peptide-78 (ENA-78) and interleukin-8
469	(IL-8) by cultured human endometrial epithelial and stromal cells. Fertil Steril
470	2008; 89 :1530–1536.
471	Bertschi D, McKinnon BD, Evers J, Bersinger NA, Mueller MD. Enhanced inflammatory
472	activity of endometriotic lesions from the rectovaginal septum. Mediators Inflamm
473	2013; 2013 :450950.
474	Bonizzi G, Karin M. The two NF-kappaB activation pathways and their role in innate and
475	adaptive immunity. Trends Immunol 2004;25:280-288.
476	Bono Y, Kyo S, Takakura M, Maida Y, Mizumoto Y, Nakamura M, Nomura K, Kiyono T,
477	Inoue M. Creation of immortalised epithelial cells from ovarian endometrioma. $Br J$
478	<i>Cancer</i> 2012; 106 :1205–1213.
479	Borrelli GM, Abrão MS, Mechsner S. Can chemokines be used as biomarkers for
480	endometriosis? A systematic review. Hum Reprod Oxf Engl 2014;29:253-266.
481	Borrelli GM, Carvalho KI, Kallas EG, Mechsner S, Baracat EC, Abrão MS. Chemokines in
482	the pathogenesis of endometriosis and infertility. J Reprod Immunol 2013;98:1-9.
483	Bruner-Tran KL, Herington JL, Duleba AJ, Taylor HS, Osteen KG. Medical management of
484	endometriosis: emerging evidence linking inflammation to disease pathophysiology.

Minerva Ginecol 2013;**65**:199–213.

- 486 Choi J, Jo M, Lee E, Kim HJ, Choi D. Differential induction of autophagy by mTOR is
- 487 associated with abnormal apoptosis in ovarian endometriotic cysts. *Mol Hum Reprod*488 2014;20:309–317.
- 489 Dai L, Gu L, Di W. MiR-199a attenuates endometrial stromal cell invasiveness through
- 490 suppression of the IKK β /NF- κ B pathway and reduced interleukin-8 expression. *Mol*
- 491 *Hum Reprod* 2012;**18**:136–145.
- 492 Dunlop EA, Tee AR. mTOR and autophagy: a dynamic relationship governed by nutrients
- 493 and energy. Semin Cell Dev Biol 2014;36:121–129.
- 494 González-Ramos R, Donnez J, Defrère S, Leclercq I, Squifflet J, Lousse J-C, Van
- 495 Langendonckt A. Nuclear factor-kappa B is constitutively activated in peritoneal
 496 endometriosis. *Mol Hum Reprod* 2007;13:503–509.
- 497 Grund EM, Kagan D, Tran CA, Zeitvogel A, Starzinski-Powitz A, Nataraja S, Palmer SS.
- 498 Tumor necrosis factor-alpha regulates inflammatory and mesenchymal responses via
- 499 mitogen-activated protein kinase kinase, p38, and nuclear factor kappaB in human
- 500 endometriotic epithelial cells. *Mol Pharmacol* 2008;**73**:1394–1404.
- Haider S, Knöfler M. Human tumour necrosis factor: physiological and pathological roles in
 placenta and endometrium. *Placenta* 2009;**30**:111–123.
- 503 Halme J, Becker S, Hammond MG, Raj MH, Raj S. Increased activation of pelvic
- 504 macrophages in infertile women with mild endometriosis. *Am J Obstet Gynecol*505 1983;145:333–337.
- 506 Harada T, Enatsu A, Mitsunari M, Nagano Y, Ito M, Tsudo T, Taniguchi F, Iwabe T,
- 507 Tanikawa M, Terakawa N. Role of cytokines in progression of endometriosis.
- 508 *Gynecol Obstet Invest* 1999;47 Suppl 1:34–39; discussion 39–40.
- 509 Hinz M, Scheidereit C. The IκB kinase complex in NF-κB regulation and beyond. *EMBO Rep*510 2014;15:46–61.
- 511 Hornung D, Ryan IP, Chao VA, Vigne JL, Schriock ED, Taylor RN. Immunolocalization and
- 512 regulation of the chemokine RANTES in human endometrial and endometriosis
- 513 tissues and cells. J Clin Endocrinol Metab 1997;82:1621–1628.

- 514 Kyo S, Nakamura M, Kiyono T, Maida Y, Kanaya T, Tanaka M, Yatabe N, Inoue M.
- 515 Successful immortalization of endometrial glandular cells with normal structural and
- 516 functional characteristics. *Am J Pathol* 2003;**163**:2259–2269.
- 517 Lam LT, Davis RE, Pierce J, Hepperle M, Xu Y, Hottelet M, Nong Y, Wen D, Adams J,
- 518 Dang L, et al. Small molecule inhibitors of IkappaB kinase are selectively toxic for
- 519 subgroups of diffuse large B-cell lymphoma defined by gene expression profiling.
- 520 Clin Cancer Res Off J Am Assoc Cancer Res 2005;11:28–40.
- 521 Lebovic DI, Chao VA, Martini JF, Taylor RN. IL-1beta induction of RANTES (regulated
- 522 upon activation, normal T cell expressed and secreted) chemokine gene expression in
- endometriotic stromal cells depends on a nuclear factor-kappaB site in the proximal
 promoter. *J Clin Endocrinol Metab* 2001;86:4759–4764.
- 525 Leconte M, Nicco C, Ngô C, Chéreau C, Chouzenoux S, Marut W, Guibourdenche J,
- 526 Arkwright S, Weill B, Chapron C, *et al.* The mTOR/AKT inhibitor temsirolimus
- 527 prevents deep infiltrating endometriosis in mice. *Am J Pathol* 2011;**179**:880–889.
- 528 Lee D-F, Kuo H-P, Chen C-T, Hsu J-M, Chou C-K, Wei Y, Sun H-L, Li L-Y, Ping B, Huang
- 529 W-C, *et al.* IKK beta suppression of TSC1 links inflammation and tumor
- angiogenesis via the mTOR pathway. *Cell* 2007;**130**:440–455.
- 531 McKinnon B, Bersinger NA, Mueller MD. Peroxisome proliferating activating receptor
- 532 gamma-independent attenuation of interleukin 6 and interleukin 8 secretion from
- 533 primary endometrial stromal cells by thiazolidinediones. *Fertil Steril* 2012;97:657–
- 534 664.
- 535 McKinnon B, Bertschi D, Wotzkow C, Bersinger NA, Evers J, Mueller MD. Glucose
- transporter expression in eutopic endometrial tissue and ectopic endometriotic
- 537 lesions. J Mol Endocrinol 2014;52:169–179.
- 538 McKinnon BD, Bertschi D, Bersinger NA, Mueller MD. Inflammation and nerve fiber
- 539 interaction in endometriotic pain. *Trends Endocrinol Metab TEM* 2015;26:1–10.

540	McKinnon BD, Kocbek V, Nirgianakis K, Bersinger NA, Mueller MD. Kinase signalling
541	pathways in endometriosis: potential targets for non-hormonal therapeutics. Hum
542	Reprod Update 2016;
543	Nap AW, Groothuis PG, Demir AY, Maas JWM, Dunselman GAJ, Goeij AFPM de, Evers
544	JLH. Tissue integrity is essential for ectopic implantation of human endometrium in
545	the chicken chorioallantoic membrane. Hum Reprod Oxf Engl 2003;18:30-34.
546	Nirgianakis K, Bersinger NA, McKinnon B, Kostov P, Imboden S, Mueller MD. Regression
547	of the inflammatory microenvironment of the peritoneal cavity in women with
548	endometriosis by GnRHa treatment. Eur J Obstet Gynecol Reprod Biol
549	2013; 170 :550–554.
550	Nisolle M, Donnez J. Peritoneal endometriosis, ovarian endometriosis, and adenomyotic
551	nodules of the rectovaginal septum are three different entities. Fertil Steril
552	1997; 68 :585–596.
553	Pahl HL. Activators and target genes of Rel/NF-kappaB transcription factors. Oncogene
554	1999; 18 :6853–6866.
555	Revised American Society for Reproductive Medicine classification of endometriosis: 1996
556	Fertil Steril 1997; 67 :817–821.
557	Samartzis EP, Samartzis N, Noske A, Fedier A, Caduff R, Dedes KJ, Fink D, Imesch P. Loss
558	of ARID1A/BAF250a-expression in endometriosis: a biomarker for risk of
559	carcinogenic transformation? Mod Pathol Off J U S Can Acad Pathol Inc
560	2012; 25 :885–892.
561	Sampson J.A. Peritoneal Endometriosis, Due to the Menstrual Dissemination of Endometrial
562	Tissue into the Peritoneal Cavity. Am J Obstet Gynecol 1928;15:101-110.
563	Santi A, Felser RS, Mueller MD, Wunder DM, McKinnon B, Bersinger NA. Increased
564	endometrial placenta growth factor (PLGF) gene expression in women with
565	successful implantation. Fertil Steril 2011;96:663-668.
566	Santulli P, Borghese B, Chouzenoux S, Vaiman D, Borderie D, Streuli I, Goffinet F, Ziegler

567 D de, Weill B, Batteux F, *et al.* Serum and peritoneal interleukin-33 levels are

568 elevated in deeply infiltrating endometriosis. *Hum Reprod Oxf Engl* 2012;27:2001–

569

2009.

- Shen F, Wang Y, Lu Y, Yuan L, Liu X, Guo S-W. Immunoreactivity of progesterone receptor
 isoform B and nuclear factor kappa-B as biomarkers for recurrence of ovarian
 endometriomas. *Am J Obstet Gynecol* 2008;199:486.e1–e486.e10.
- 573 Somigliana E, Infantino M, Candiani M, Vignali M, Chiodini A, Busacca M, Vignali M.
- 574 Association rate between deep peritoneal endometriosis and other forms of the
- 575 disease: pathogenetic implications. *Hum Reprod Oxf Engl* 2004;19:168–171.
- 576 Stickles XB, Marchion DC, Bicaku E, Al Sawah E, Abbasi F, Xiong Y, Bou Zgheib N, Boac
- 577 BM, Orr BC, Judson PL, *et al.* BAD-mediated apoptotic pathway is associated with
 578 human cancer development. *Int J Mol Med* 2015;**35**:1081–1087.
- 579 Yan J, Xiang J, Lin Y, Ma J, Zhang J, Zhang H, Sun J, Danial NN, Liu J, Lin A. Inactivation
- 580 of BAD by IKK inhibits TNFα-induced apoptosis independently of NF-κB activation.
 581 *Cell* 2013;152:304–315.
- 582 Young VJ, Brown JK, Saunders PTK, Duncan WC, Horne AW. The peritoneum is both a
- 583 source and target of TGF-β in women with endometriosis. *PloS One* 2014;9:e106773.
- 584 Zeitvogel A, Baumann R, Starzinski-Powitz A. Identification of an invasive, N-cadherin-
- 585 expressing epithelial cell type in endometriosis using a new cell culture model. *Am J*
- 586 Pathol 2001;159:1839–1852.
- 587
- 588

589 Table 1; Comparison of endometriotic lesion and peritoneal fluid protein expression in

590 samples removed from women using different hormonal treatment. Data are Mean ±

591 SEM

	Hormone treatment					
	No hormone	Combined oral contraceptive	GnRH analogue	Total		
	(n=14)	(n = 4)	(n=3)	(n =21)	р	
ρΙΚΚα/β (%)	66 ± 9.8	77 ± 27.8	92 ± 17.5	72 ± 8.5	0.4732	
ΙΚΚβ (%)	120 ± 23.5	90 ±27.7	151 ± 27.7	119 ± 16.9	0.3819	
TNFα (pg/ml)	2.1 ± 0.61	1.2 ± 0.53	1.5 ± 0.54	1.9± 0.43	0.7721	

592 -IKK β and pIKK α/β protein values were determined via Semi-quantitative Western blot 593 densitometry analysis and expressed as a percentage of a standardised control sample 594 included in each Western blot, as described in detail in the methods section.

595 -TNF α values represent peritoneal fluid TNF α concentrations determined by ELISA and 596 expressed as pg/ml.

597 -Analysis of significance was performed via the non-parametric One-way ANOVA test

598 (Kruskal-Wallis) and a post hoc comparison of all groups (Dunn's multiple comparison test)

significance p < .05

600

602 Table II: Comparison of protein expression in formalin fixed paraffin embedded

603 endometriotic tissue according to hormonal treatment. Data are mean \pm SEM

Hormonal use	No hormone	Combined oral contraceptive	GnRH analogue	Total	
	(n= 6)	(n = 5)	(n=5)	(n =16)	*P
ρΙΚΚα/β					
Epithelial	1.7 ± 0.56	3.4 ± 1.78	6.2 ± 2.27	3.6 ± 0.98	0.108
Stromal	0.3 ± 0.33	0.8 ± 0.37	0.6 ± 0.60	0.5 ± 0.24	0.448
ΙΚΚβ					
Epithelial	2.3 ± 0.95	2.2 ± 1.11	3.0 ± 1.76	2.5 ± 0.70	0.949
Stromal	0.7 ± 0.42	12.4 ± 1.75	0.0 ± 0.00	1.00 ± 0.58	0.327

604

-Values represent the scores derived from the semi-quantitative IHC analysis, described in 605 detail in the methods.

606 -Analysis of significance was performed via the non-parametric One-way ANOVA test

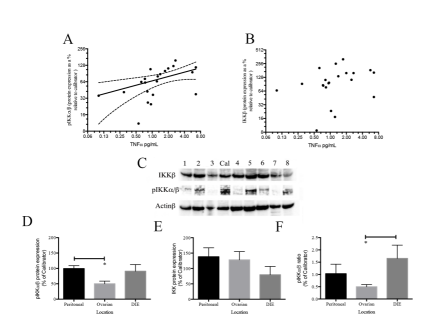
607 (Kruskal-Wallis) and a post hoc comparison of all groups (Dunn's multiple comparison test)

608 significance p < .05

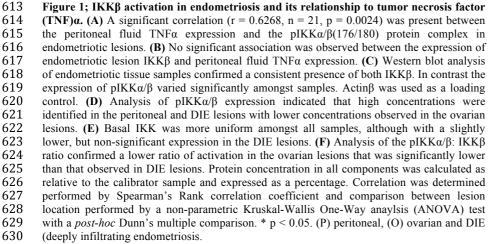
609

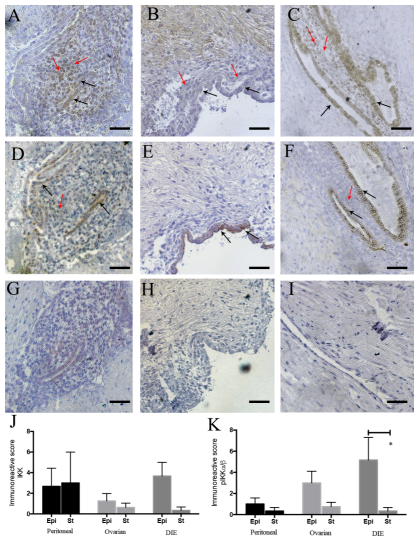
Brett McKinnon 7.2.2017 09:39

Kommentar [7]: A typo resulted in an additional 1 at the start of the number. The correct value is 2.4

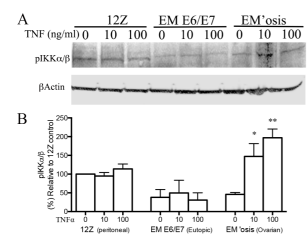






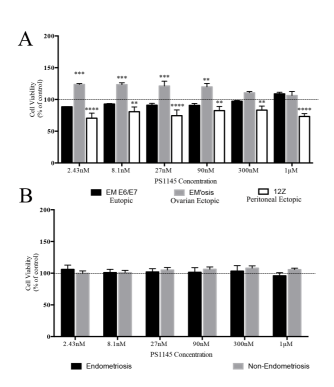


632 633 Figure 2; Cell specific expression and activation of IKKß in endometriotic lesions. Basal 634 IKKβ expression was observed in both epithelial (black arrows) and stromal cells (red 635 arrows) of endometriotic lesions removed from the (A) peritoneal, (B) ovarian, and (C) DIE 636 regions. The expression of the activated IKK complex (pIKK α/β) was limited predominantly 637 to the epithelial cells (black arrows), although some stromal cell expression was observed 638 (red arrows). This was consistent across lesions from the (D) peritoneal, (E) ovarian and (F) 639 and DIE lesions. Negative controls showed no expression in lesions from the (G) peritoneal 640 (H) ovarian, or (I) DIE region. A semi-quantitative analysis of the cell specific expression 641 indicated that no statistically significant variation in (J) IKKB expression was observed 642 between epithelial and stromal cells, however pIKK α/β was significantly stronger in the 643 epithelial cells for all lesions with a largest difference observed in the DIE lesions. (K) 644 Analysis of protein activation and expression between cell type and lesion location was 645 performed with a Two-way analysis of variance (ANOVA) test with a post-hoc Tukey's 646 multiple comparison. Scale bars = $50\mu m$. * p < 0.05 647



649 650 651 Figure 3; pIKKa/ β after tumor necrosis factor (TNF)a treatment in endometriotic epithelial cells. (A) Western blot analysis confirmed the expression of $pIKK\alpha/\beta$ in the 12Z, 652 EM E6/E7 and EM 'osis cell lines both with and without $TNF\alpha$ treatment. (B) Semi-653 654 quantitative analysis indicated TNF α did not significantly influence either 12Z, or EM E6/E7 expression. There was however a significant increase in $pIKK\alpha/\beta$ after both 10ng/ml and 100ng/ml TNFα in the EM'osis cell line.

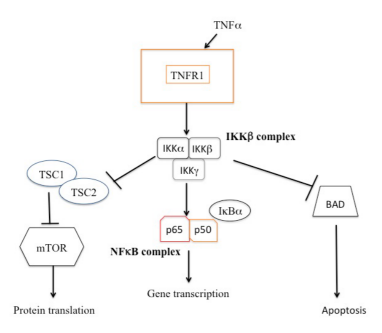
655 656



⁶⁵⁸ 659

657

Figure 4; Influence of IKKB inhibition on epithelial and stromal cell viability. (A) 660 Inhibition of IKKB activity with increasing concentrations of PS1145 had no influence on the 661 eutopic derived epithelial EM E6'E7 cells, significantly increased the cell viability of the 662 ovarian derived EM'osis cells, significantly reduced the viability of the peritoneal derived 663 12Z cells. (B) PS1145 had no influence on stromal cells from women with and without 664 endometriosis. Analysis on the influence between cell type and PS1145 concentrations on cell viability performed with a Two-way analysis of variance (ANOVA) test with a post-hoc Tukey's multiple comparison test. ** p < 0.01, *** p < 0.001, **** p < 0.0001. 665 666





668 669 Figure 5; The IKK complex and its influence on downstream signalling pathways. 670 Binding of extracellular tumor necrosis factor (TNF)a to TNF receptor 1 (TNFR1) stimulates 671 the phosphorylation and activation of IKK α and IKK β that exist as a complex along with 672 IKKγ. Activation of IKKβ leads to the phosphorylation of IκBα, which under basal conditions 673 is bound to the p65 subunit of the NF κB complex. Phosphorylation of I $\kappa B \alpha$ removes it from 674 675 the NF κ B complex and initiates protesomal degradation allowing NF κ B translocation into the nucleus and gene transcription. Activation of the IKK complex can also lead to an interaction 676 with the tuberous sclerosis (TSC)2 protein that exists in a heterodimer with TSC1. Inhibition 677 of TSC2 activity increases the activity of the mammalian target of rapamycin (mTOR) 678 complex stimulating both protein translation and cellular proliferation. Activation of the IKK 679 complex also leads to an inactivation of the BH3 only BAD protein inactivating TNFa 680 stimulated apoptosis. 681