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2	VICEVERSA, BECAUSE OF D	RUG INTOLERANCE: DOES TH	IE CHANGE BENEFIT
3	WOMEN WITH ENDOMETRIC	OSIS?	
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23	SHORT TITLE: Intolerance to early a statement of the stat	strogen-progestins and progestins f	or endometriosis
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SHIFTING FROM ORAL CONTRACEPTIVES TO NORETHISTERONE ACETATE, OR

27 ABSTRACT

28 Background/Aims: Oral contraceptives (OC) and norethisterone acetate (NETA) are among first-

29 line medical therapies for symptomatic endometriosis, but their use is sometimes associated with

30 intolerable side effects. We investigated whether shifting from low-dose OC to NETA (2.5 mg/day),

31 or viceversa, improved tolerability.

32 Methods

33 Sixty-seven women willing to discontinue their treatment because of intolerable side effects despite

34 good pain relief, were enrolled in a self-controlled study, and shifted from OC to NETA (n = 35) or

from NETA to OC (n = 32). The main study outcome was satisfaction with treatment 12 months

36 after the change. Tolerability, pain symptoms, health-related quality of life, psychological status,

37 and sexual functioning were also evaluated.

38 **Results**

39 After treatment change, good tolerability was reported by 37% of participants who shifted to

40 NETA, and by 52% of those who shifted to OC. At 12-month assessment, 51% of women intolerant

41 to OC were satisfied with NETA, and 65% of those intolerant to NETA were satisfied with OC

42 (intention-to-treat analysis). Other study variables did not vary substantially.

43 Conclusions

44 In selected endometriosis patients, shifting from OC to NETA, or viceversa, because of side effects,

45 improved tolerability. Better results were observed when substituting NETA with OC rather than

the other way round.

47 INTRODUCTION

48 Combined oral contraceptives (OC) and progestins are indicated by major international guidelines 49 as the first-line medical treatment options for women not seeking conception and with 50 endometriosis-associated pelvic pain [1-4]. Overall, about two thirds of patients appear to benefit 51 from these therapies [5-16]. The main reason of treatment failure in the remaining third, in addition 52 to inefficacy, is drug intolerance. As untoward effects of OCs and progestins partly differ, a shift 53 from the former to the latter compounds, or viceversa, could allow continuing treatment with a safe, 54 effective, and unexpensive medication without the need for stepping up to a drug with a less favorable therapeutic profile or resorting to surgery. However, very limited information is available 55 56 on what should a patient expect from these changes [17, 18]. The answers to these questions seem 57 important as the clinical issue is not rare and may interfere with health-related quality of life and 58 disease management. Given this background, we sought to investigate whether shifting from an OC 59 to a progestin, or viceversa, specifically because of drug intolerance, is of benefit in terms of relief 60 from side effects and, in case these measures are effective, whether they imply reduced efficacy on 61 pain symptoms.

62 MATERIALS AND METHODS

The main objective of the present study was to assess the proportion of patients satisfied with their therapy 12 months after a change from a low-dose, monophasic OC to norethisterone acetate (NETA), or viceversa, because of side effects intolerable to the point of requesting treatment discontinuation. Therefore, in the present study population, patient dissatisfaction was not caused by inefficacy on pain symptoms. Secondary objective was the evaluation of variations in pain symptoms, health-related quality of life, psychological status, and sexual function associated with the shift from OC to NETA, or viceversa.

A prospective, self-controlled study design was adopted because it allows within-person
 comparisons avoiding the potential confounding caused by differences between patients [19]. The
 investigation was performed in an academic department specializing in the management of

endometriosis, and the competent Institutional Review Board approved the study. Patients signed an

74 informed consent form before enrollment. Women who denied their consensus were excluded.

75 Patients

We considered 18- to 40-year old women, not seeking conception, with a surgical diagnosis of endometriosis in the previous 24 months or with a current non-surgical diagnosis of endometriosis, and using an OC or NETA for pelvic pain, but unwilling to continue the current treatment because of dissatisfaction due to intolerable side effects. Non-surgical diagnoses were based on previously published criteria [20-22]. Participants were recruited during the period August 2014 - July 2015.

82 Women were informed that: i) OC or NETA may, in some women, cause side effects, 83 frequently because of the estrogen component in the former case, and of residual androgenic 84 activity in the latter case; ii) switching to, respectively, a progestin monotherapy or an OC 85 containing another type of progestin could result in subjective improvement; iii) also the alternative 86 drug was associated with side effects, and the efficacy of the proposed change of therapy was 87 uncertain; iv) OCs and progestins are indicated by major international guidelines as the first-line 88 treatment for endometriosis-associated pelvic pain [1-4], but that other medical therapies exist, 89 although characterized by a less favorable balance between benefits, harms and costs [23-27]; v) 90 laparoscopic surgery was a reasonable alternative in case they declined a change in pharmacological 91 treatment, but that pain and lesion recurrence was about 10% a year without long-term 92 postoperative medical therapy [28, 29].

93 Treatments

94 <u>a. Switch from OC to NETA</u>

95 Norethisterone acetate, a 19-nortestosterone derivative progestin, has been repeatedly evaluated in 96 women with endometriosis [6, 9-11, 30-32], and has been routinely used in our referral center for 97 several years [7, 14-16]. Norethisterone acetate is approved by the FDA and the Italian Ministry of 98 Health for the treatment of endometriosis and is reimbursed by the Italian National Health System.

99	Norethisterone acetate was prescribed at the dose of 2.5 mg once a day, per os. The progestin was
100	started after 4-7 days since OC discontinuation, depending on the type of OC previously used.
101	b. Switch from NETA to OC
102	The OCs used in our center were monophasic formulations containing ethinyl-estradiol 0.015 mg
103	and gestodene 60 mg or, in case of spotting, ethinyl-estradiol 0.02 mg and desogestrel 150 mg. In
104	smokers and in those with a BMI \ge 30, a combination of ethinyl-estradiol 0.02 mg and
105	levonorgestrel 100 mg was prescribed. Women were allowed to choose between cyclic and
106	continuous OC use based on their preference because the reason for the change of medication was
107	intolerance, not inefficacy on pain. A pause without treatment was not suggested before starting
108	OC.
109	NETA and OC were continued without preplanned time limits. However, for the purpose of
110	the present study, only the first 12 months of use have been evaluated. In case of prolonged spotting
111	$(\geq 7 \text{ days})$ or breakthrough bleeding during NETA or continuous OC use, the patients were advised
112	to discontinue treatment for one week in the former case, and 4-7 days in the latter case.
113	Measurements
114	All patients assisted in our center systematically undergo clinical and ultrasonographic
115	evaluation every six months. On these occasions, women are routinely asked to complete five
116	questionnaires, two on pain (a numeric rating scale, NRS; and a multi-dimensional categorical
117	rating scale, MCRS), one on quality of life (the Short Form-12 questionnaire, SF-12), one on
118	psychological status (the Hospital Anxiety and Depression scale, HADS), and one on sexual
119	functioning (the Female Sexual Function Index, FSFI). Women are also asked to indicate drug
120	tolerability using a NRS and to rate the degree of satisfaction with their treatment.
121	The above scales and questionnaires have been described previously in detail [7, 13-16]. The
122	presence and severity of dysmenorrhea, deep dyspareunia, non-menstrual pelvic pain, and dyschezia
123	were assessed using an 11-point NRS, with 0 indicating absence of pain and 10 pain as bad as it
124	could be. Patients were also asked to grade the severity of the above symptoms using a 0- to 3-point

125 MCRS modified from that devised by Biberoglu and Behrman [33]. Irregular bleeding during 126 treatment was defined as spotting (scanty bleeding requiring < 1 pad or tampon per day) or 127 breakthrough bleeding (light or moderate bleeding requiring ≥ 2 pads or tampons per day). Pain 128 during spotting or breakthrough bleeding was considered as dysmenorrhea. 129 The SF-12 health survey, developed from the original SF-36 questionnaire [34, 35], is a well 130 know, validated self-administered 12-item instrument. It measures health dimensions covering 131 functional status, well-being, and overall health. Information from the 12 items is used to construct 132 physical (PCS-12) and mental (MCS-12) component summary measures [36, 37], with higher scores indicating better health perception. 133 134 The HADS questionnaire is a self-assessment mood scale specifically designed for use in 135 non-psychiatric hospital outpatients to determine states of anxiety and depression. It comprises 14 136 questions, 7 for the anxiety subscale and 7 for the depression subscale. Lower scores indicate better 137 psychological status [38]. 138 The FSFI questionnaire is a 19-item, multidimensional, self-report instrument for evaluating 139 the main categories of female sexual dysfunction and sexual satisfaction [39-41]. The transformed 140 maximum score for each domain is 6, and the maximum (best) transformed full-scale score is 36, 141 with a minimum full-scale score of 2.0. 142 Occurrence of side effects associated with medical treatments is actively investigated in our 143 endometriosis outpatient clinic, and the overall tolerability of hormonal therapies is measured using 144 a 0- to10-point NRS, with 0 indicating absolutely intolerable untoward effects and 10 absence of 145 adverse effects. Scores are then categorized, with 9-10 indicating that a drug is very well tolerated; 7-8, well tolerated; 5-6, moderately tolerated; 3-4, poorly tolerated; 0-2, not tolerated [16]. 146 147 Patients rated the degree of satisfaction after the modification of their treatment according to 148 a five-category scale (very satisfied, satisfied, neither satisfied nor dissatisfied, dissatisfied, very 149 dissatisfied) by answering the following question: "Taking into consideration the variations

150 occurred in side effects and overall tolerability of treatment, pain symptoms, physical and

151 psychological well-being, health-related quality of life, and sexual functioning, how would you

152 define the level of satisfaction with your current treatment?" In order to limit the potential effect of

153 confounding, satisfaction with treatment, the main study outcome, was dichotomized into

154 "satisfied" (very satisfied plus satisfied) and "dissatisfied" (neither satisfied nor dissatisfied plus

155 dissatisfied plus very dissatisfied).

156 Data management

157 The focus of the investigation was not a head-to-head comparison between OC and NETA but,

158 instead, quantification of the proportion of women who were satisfied with a change in treatment 12

159 months after OC or NETA discontinuation because of intolerance. No study is available to define

160 the potential benefits of shifting from OC to NETA or viceversa in this clinical condition.

161 Therefore, a pre-planned power calculation was not performed, and we decided to include all the

162 eligible patients evaluated in a 12-month period.

163 Data were archived using Excel 2003 (Microsoft Corporation, Redmond, Washington, U.S.A.) and exported in SPSS 18.0 (SPSS, Inc, Chicago, IL, U.S.A.) or SAS software 9.4 (SF-12 164 165 data; SAS Institute Inc., Cary, NC, U.S.A.) for statistical analysis. Estimate of patient satisfaction 166 rate was performed according to the intention-to-treat principle, considering as dissatisfied all patients who dropped out of the study for any reason except conception seeking, thus including 167 168 request for surgery and lost to follow-up. Variations in drug tolerability, pelvic pain symptoms, health-related quality of life, psychological status, and sexual functioning between baseline and 12-169 month values were evaluated by using the paired Student t test for normally distributed data, the 170 171 non-parametric Wilcoxon matched pairs test for non-normally distributed data, the McNemar test for categorical variables, and the Fisher Exact test in case of cells without numerical data. 172 173 Determinants of satisfaction with treatment were investigated with unpaired tests (Student t test for 174 normally distributed continuous variables, Wilcoxon test for non-normally distributed continuous variables, and the chi-squared test for categorical variables). All statistical tests were two-sided. A P 175

176 value < 5% was considered statistically significant. When appropriate, 95% confidence intervals

177 (CIs) were calculated for the observed differences by applying a binomial distribution model.

178 RESULTS

179 A total of 35 women shifted from OC to NETA, and 32 from NETA to OC. The distribution of

180 demographic and clinical characteristics of the patients in the two study groups are shown in Table

181 1.

182 a. Switch from OC to NETA

183 The median duration [interquartile range, IQR] of OC use was 6 months [3-14]. Nineteen women 184 (54%) were using OC cyclically and 16 (46%) continuously. The most frequent untoward effects 185 that determined the request for OC discontinuation despite an appreciable effect on pain symptoms 186 were headache (49%), breakthrough bleeding (14%), and weight gain (11%). Eight women (23%) 187 dropped out from the study between the 6- and 12-month evaluation owing to persistence of 188 (headache, n = 3) or onset of different (mood changes, n = 1; urticarial rash, n = 1; breakthrough 189 bleeding, n = 1) side effects, onset of non-menstrual pelvic pain (n = 1), and unwillingness to 190 undertake any further treatment (n = 1). Variation of frequency of side effects associated with the 191 shift from OC to NETA in the 27 women who completed the 12-month study period is reported in 192 Table 2. None of the differences were statistically significant. A trend was observed toward a 193 decrease in frequency of headache (from 56% to 30%) and an increase in that of weight gain (from 194 30% to 44%). However, the severity of untoward effects decreased significantly, as the mean \pm SD 195 tolerability NRS score increased from 3.0 ± 1.6 to 5.7 ± 2.4 (P < 0.001). Ten women (37%) 196 reported good or very good (NRS \geq 7) drug tolerability, compared with none at baseline. 197 The severity of pain symptoms did not vary significantly except for dysmenorrhea that 198 decreased at evaluation by means of the NRS (Table 3). Overall, the frequency of moderate or 199 severe complaints was marginal at both baseline and 12-month assessment. No substantial 200 variations were observed also in psychological status and sexual functioning. With regard to health-201 related quality of life, a significant improvement was reported only in the physical component of the

202	SF-12 questionnaire (Table 3). At the end of the study period 18/35 (51%; 95% C.I., 36% to 67%)
203	women were satisfied or very satisfied with the treatment change, whereas $17/35$ (49%; 95% C.I.,
204	33% to 64%) were neither satisfied nor dissatisfied, dissatisfied, or very dissatisfied. All the patients
205	who dropped out from the study were included as dissatisfied in this intention-to-treat analysis.
206	b. Switch from NETA to OC
207	The median duration [interquartile range, IQR] of NETA use was 12 months [5-22]. The most
208	frequent untoward effects that determined the request for NETA discontinuation despite an
209	appreciable effect on pain symptoms were weight gain (19%), headache (16%), breakthrough
210	bleeding (16%), decreased libido (16%), spotting (12%), and mood changes (12%). Seven women
211	(22%) dropped out from the study between the 6- and 12-month evaluation owing to persistence of
212	(acne, $n = 1$) or onset of different (headache, $n = 4$) side effects, onset of non-menstrual pelvic pain
213	(n = 1), and pregnancy desire $(n = 1)$. Variation of frequency of side effects associated with the shift
214	from NETA to OC in the 25 women who completed the 12-month study period is reported in Table
215	4. Again, none of the differences were statistically significant. A trend was observed toward a
216	decrease in frequency of weight gain (from 36% to 16%), decreased libido (from 44% to 32%), and
217	vaginal dryness (from 36% to 24%), and an increase in that of headache (from 20% to 40%).
218	However, the severity of untoward effects decreased significantly, as the mean \pm SD tolerability
219	NRS score increased from 3.5 ± 1.7 to 6.9 ± 2.5 ($P < 0.001$). Thirteen women (52%) reported good
220	or very good (NRS \geq 7) drug tolerability, compared with none at baseline.
221	Based on NRS assessment, the severity of deep dyspareunia and non-menstrual pelvic pain
222	decreased significantly (Table 5). A trend was observed toward a decrease in the frequency of
223	moderate to severe deep dyspareunia (eight women at baseline vs three at 12 months) and dyschezia
224	(four and two women, respectively) at MCRS evaluation (Table 5). Significant improvements were
225	observed in both the anxiety and depression HADS subscales scores, as well as in the FSFI scores.
226	No significant variations were reported in both the physical and the mental components of the SF-
227	12 questionnaire (Table 5). One woman who dropped out of the study because of pregnancy desire

228 was not considered in the intention-to-treat analysis of satisfaction with treatment. At the end of the

study period 20/31 (65%; 95% C.I., 47% to 79%) patients were satisfied or very satisfied with the

treatment change, whereas 11/31 (35%; 95% C.I., 20% to 52%) were neither satisfied nor

231 dissatisfied, dissatisfied, or very dissatisfied.

232 DISCUSSION

Overall, the main finding of the present study was that, when OC or NETA are not tolerated, shifting to the other compound allows the majority of patients with endometriosis to improve tolerability and to continue medical treatment with a safe, effective, and unexpensive drug. The benefit seems larger when the shift is from NETA to OC rather than the other way round, as the proportion of satisfied patients at the end of the study period was, respectively, 65% and 51%. Moreover, in the latter case the 95% C.I.s of the rates of satisfied and dissatisfied women amply overlapped, whereas in the former case the 95% C.I. overlapping was marginal.

Considering a shift from OC to NETA may be beneficial especially in women experiencing headache, as previously suggested by Morotti *et al.* [17]. The frequency of the other untoward effects associated with OC use were not reduced, but their severity was, as demonstrated by the increase in 12-month follow-up NRS tolerability score compared with baseline values.

244 Considering a shift from NETA to OC may be beneficial especially in women experiencing 245 side effects typically associated with this type of progestin, such as weight gain, acne, bloating, and 246 decreased libido. On the other hand, this change may lead to an increase in the frequency of 247 headache, likely associated with the estrogen component. This confirms that OCs with the lowest 248 possible estrogen dose should be chosen also in women with endometriosis in order to improve both 249 safety and tolerability [42-45].

The larger effect observed when the shift was from NETA to OC confirms that low-dose, monophasic estrogen-progestin combinations should retain their role in the management of endometriosis, provided pain symptoms are adequately relieved. In this regard, it should be highlighted that at baseline pain was generally well controlled in both study groups, and that the focus here was on tolerability, not efficacy on symptoms. This also explains the limited significant variations in pain symptoms' severity independently of the direction of the change between the two medications, demonstrating that the observed amelioration of tolerability was not at detriment of efficacy on pain. Conversely, marginal improvements in the severity of dysmenorrhea when shifting from OC to NETA, and of deep dyspareunia and non-menstrual pain when shifting from NETA to OC were reported, although of questionable clinical importance.

260 Our study has limitations. The combination of the observational design with the limited 261 sample size increases the risk of confounding. Moreover, the population was highly selected, and this precludes generalization of the results to endometriosis patients with different complaint types. 262 263 However, the self-controlled design was chosen purposely because the objective of the study was to 264 assess variations in tolerability when shifting to NETA or OC not in a general population using the 265 other drug, but specifically in those patients who were dissatisfied because of intolerable side 266 effects and that would have otherwise discontinued medical therapy. In a self-control study, 267 recruited patients act as their own control, thus limiting the effect of confounding. In fact, study 268 outcomes may be influenced by relevant characteristics that may differ between patients [19]. In 269 addition, overoptimistic results should have been avoided, as patient satisfaction was assessed 270 including all dropouts as dissatisfied.

271 The period of use of OC and NETA before changing medication was fairly long. Thus, the phenomenon of regression toward the mean seems unlikely, given that the clinical condition was 272 273 chronic and that all study variables were measured repeatedly before enrollment. Also a carry-over 274 effect should be ruled out, as the baseline patients' conditions were the worst possible in terms of tolerability. Therefore, if a carry-over effect was in play, this was detrimental, not beneficial, again 275 276 potentially leading to conservative estimates. Also a placebo effect cannot be excluded. However, given the long study period, this seems little probable, as the placebo effect may not last for one 277 278 year when drug tolerability is unacceptable.

279 The proportion of dropouts was high and above the usually indicated 20% cut-off over 280 which the study findings are considered of questionable validity [46]. However, this cut-off may not 281 be appropriate when all patients at recruitment are considering abandonment of medical treatment 282 owing to dissatisfaction. In these conditions, a 22-23% dropout rate may even appear fairly low. 283 Owing to the limited number of participants, the analysis of determinants of success was 284 deemed unreasonable. More in general, the small sample size could have led to some type II errors. 285 thus impeding the identification of potential factors predictive of satisfaction with treatment change. 286 On the other hand, in our experience it is not easy for endometriosis patients to decide to 287 discontinue a medical therapy that is effective on pain, solely because of side effects. In this regard, 288 it may not be excluded that women referred or self-referred to our center are more motivated to 289 choose medical rather that surgical treatment. If this was true, such selection bias would render 290 generalization of the study results more problematic.

However, when discussing generalization, we also believe that our findings provide a realistic picture of what happens in everyday practice, and our data may help clinicians when counselling patients experiencing upsetting untoward effects with OC or NETA. Observational studies may be very helpful in assessing the real world effectiveness of treatments that have already been demonstrated to work in highly controlled research settings [47], as OCs and NETA in women with symptomatic endometriosis [5, 7, 8, 12].

It could also be argued that, in women who were intolerant to NETA, instead of suggesting OC we could have suggested shifting to dienogest, that has been proven to be better tolerated than NETA [16]. However, many women assisted in our center cannot afford the cost of dienogest (\notin 730 - $\$860 - \pounds 670$ per year in Italy, not reimbursed by the Italian NHS) and prefer NETA (\pounds 18 - \$21 - \pounds 17 per year in Italy, \pounds 4 per year when reimbursed by the Italian NHS) specifically for economic reasons. Indeed, we previously demonstrated that the cost of dienogest limited its effectiveness despite its good tolerability [16]. Moreover, here the issue was not poor pain control, but drug

304	intolerance, and indeed the larger benefit was observed precisely when shifting from NETA to OC.
305	Thus, changing for dienogest would have led to waste of money in the majority of patients.
306	In conclusion, when endometriosis-associated pain was relieved by OC or NETA, but the
307	medications could no longer be used because of intolerable side effects, shifting to the other
308	compound resulted in substantial improvement of tolerability in the majority of women. The change
309	of therapy was particularly beneficial in patients using NETA who shifted to OC. Women should be
310	informed about this further therapeutic option in order to be enabled to choose a treatment
311	modification that is aligned with their preferences and priorities.
312	

313 DISCLOSURE STATEMENT (CONFLICT OF INTEREST)

PV, FO, MPF, LB, and AR declare that they have no conflicts of interest. ES received grants from
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444	

Characteristic	From OC to NETA study group (n = 35)	From NETA to OC study group $(n = 32)$
Age (years)	35.5 ± 4.7	34.2 ± 5.3
BMI (Kg/m ²)	23.6 ± 4.0	20.5 ± 2.6
Smoking	6 (17%)	9 (28%)
Previous deliveries	15 (43%)	6 (19%)
Previous surgical procedures for endometriosis		
None	9 (26%)	14 (44%)
1	18 (51%)	11 (34%)
2	7 (20%)	5 (16%)
\geq 3	1 (3%)	2 (6%)
Endometriotic lesion type ^a		
Deep infiltrating endometriosis	17 (49%)	24 (75%)
Ovarian endometriomas	28 (80%)	18 (56%)
Pain symptoms ^b		
Dysmenorrhea	15 (42%)	8 (25%)
Deep dyspareunia	5 (15%)°	15 (30%) ^d
Non-menstrual pelvic pain	5 (14%)	12 (37%)
Dyschezia	2 (6%)	7 (22%)
Duration of previous treatment [months]	6 [3-14]	12 [5-22]

Table 1. Distribution of baseline demographic and clinical characteristics of women who shifted to norethisterone acetate (NETA) for intolerance to low-dose oral contraceptive (OC), and of women who shifted to OC for intolerance to NETA.

Data is reported as mean ± SD, or number (percentage), or median [interquartile range].

NETA = norethisterone acetate. OC = low-dose, combined oral contraceptive. BMI = body mass index.

^a The sum does not add to the total as some women had both lesion types.

⁴⁴⁶ ^bNumeric rating scale > 0. Mild pain symptoms are also included.

447 °One woman did not have sexual intercourses at basal and/or at 12-month evaluation.

448 ^dTwo women did not have sexual intercourses at basal and/or at 12-month evaluation

Side effect ^a	Baseline evaluation	12-month evaluation	Р
Headache	15 (56%)	8 (30%)	NS
Spotting	5 (18%)	6 (22%)	NS
Breakthrough bleeding	1 (4%)	0 (0%)	NS
Weight gain	8 (30%)	12 (44%)	NS
Nausea	2 (7%)	1 (4%)	NS
Decreased libido	7 (26%)	5 (18%)	NS
Vaginal Dryness	4 (15%)	6 (22%)	NS
Bloating or swelling	5 (18%)	6 (22%)	NS
Breast tenderness	0 (0%)	4 (15%)	NS
Acne	0 (0%)	3 (11%)	NS
Alopecia	0 (0%)	0 (0%)	NS
Mood changes	5 (18%)	5 (18%)	NS
Others	11 (41%)	9 (33%)	NS

Table 2. Per-protocol analysis of frequency of side effects reported at baseline and at 12-month evaluation by patients (n = 27) shifting from OC to NETA.

^aSome women reported more than one side effect.

Data are number (percentage).

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Table 3. Per-protocol analysis^a of pain symptoms, health-related quality of life, psychological status, and sexual functioning scores variation between baseline and 12month evaluation in patients (n = 27) shifting from OC to NETA.

Symptoms / Questionnaires	Baseline evaluation	12-month evaluation	Р
Dysmenorrhea			
NRS	0 [0-4]	0 [0-0]	0.01
MCRS ≥ 2	2 (7%)	0 (0%)	NS
Deep dyspareunia ^b			
NRS	0 [0-0]	0 [0-0]	NS
MCRS ≥ 2	1 (4%)	2 (8%)	NS
Non-menstrual pelvic pain			
NRS	0 [0-0]	0 [0-0]	NS
MCRS ≥ 2	1 (4%)	0 (0%)	NS
Dyschezia			
NRS	0 [0-0]	0 [0-0]	NS
MCRS ≥ 2	1 (4%)	0 (0%)	NS
SF-12			
Physical component	50.0 ± 11.1	55.4 ± 4.5	0.03
Mental component	40.0 ± 11.7	42.6 ± 13.2	NS
HADS			
Anxiety	6.6 ± 4.3	5.9 ± 4.6	NS
Depression	5.8 ± 4.3	5.4 ± 5.1	NS
Total	12.4 ± 8.1	11.3 ± 9.1	NS
FSFI total score ^b	26.2 ± 5.7	26.2 ± 6.7	NS

Data is reported as mean ± SD, or number (percentage), or median [interquartile range]. NRS = 0 to 10-point numeric rating scale. MCRS = 0 to 3-point multidimensional categorical rating scale modified from that devised by Biberoglu and Behrman [33]. SF-12 = Short Form-12 [36, 37]. HADS = Hospital Anxiety and Depression Scale [38]. FSFI = Female Sexual Function Index [39, 40].

NS = not significant.

^aWomen who withdrew before 12-month follow-up assessment (n = 8) were excluded. ^bOne woman did not have sexual intercourses either at baseline and/or at 12-month evaluation.

Side effect ^a	Baseline evaluation	12-month evaluation	Р
Headache	5 (20%)	10 (40%)	NS
Spotting	4 (16%)	7 (28%)	NS
Breakthrough bleeding	3 (12%)	0 (0%)	NS
Weight gain	9 (36%)	4 (16%)	NS
Nausea	2 (8%)	0 (0%)	NS
Decreased libido	11 (44%)	8 (32%)	NS
Vaginal Dryness	9 (36%)	6 (24%)	NS
Bloating or swelling	4 (16%)	2 (8%)	NS
Breast tenderness	0 (0%)	0 (0%)	NS
Acne	2 (8%)	0 (0%)	NS
Alopecia	0 (0%)	0 (0%)	NS
Mood changes	5 (20%)	1 (4%)	NS
Others	5 (20%)	1 (4%)	NS

Table 4. Per-protocol analysis of frequency of side effects reported at baseline and at 12-month evaluation by patients (n = 25) shifting from NETA to OC.

^aSome women reported more than one side effect.

Data are number (percentage).

Symptoms / Questionnaires	Baseline evaluation	12-month evaluation	Р
Dysmenorrhea			
NRS	0 [0-1.5]	0 [0-3]	NS
MCRS ≥ 2	0 (0%)	1 (4%)	NS
Deep dyspareunia ^b			
NRS	5 [0-8]	0 [0-5.5]	0.02
MCRS ≥ 2	8 (35%)	3 (13%)	NS
Non-menstrual pelvic pain			
NRS	0 [0-4.5]	0 [0-0]	0.02
MCRS ≥ 2	2 (8%)	1 (4%)	NS
Dyschezia			
NRS	0 [0-1.5]	0 [0-0]	NS
MCRS ≥ 2	4 (16%)	2 (8%)	NS
HADS			
Anxiety	4.7 ± 3.5	3.6 ± 3.2	0.02
Depression	5.4 ± 4.0	3.8 ± 3.4	0.03
Total	10.1 ± 7.3	7.4 ± 6.3	0.02
SF-12			
Physical component	52.8 ± 9.1	54.8 ± 4.4	NS
Mental component	42.1 ± 11.7	46.1 ± 10.0	NS
FSFI total score ^b	21.9 ± 8.6	25.4 ± 7.9	0.01

Table 5. Per-protocol analysis^a of pain symptoms, health-related quality of life, psychological status, and sexual functioning scores variation between baseline and 12-month evaluation in patients (n = 25) shifting from NETA to OC.

Data is reported as mean \pm SD, or number (percentage), or median [interquartile range]. NRS = 0 to 10-point numeric rating scale. MCRS = 0 to 3-point multidimensional categorical rating scale modified from that devised by Biberoglu and Behrman [33].

SF-12 = Short Form-12 [36, 37]. HADS = Hospital Anxiety and Depression Scale [38].

FSFI = Female Sexual Function Index [39, 40].

NS = not significant.

^aWomen who withdrew before 12-month follow-up assessment (n = 7) were excluded.

^bOne woman did not have sexual intercourses either at baseline and/or at 12-month evaluation.