Can a bothersome course of pelvic pain from mid-pregnancy to birth be predicted? A Norwegian prospective longitudinal SMS-Track study

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ABSTRACT

Objective To explore if pregnant women with pelvic girdle pain (PGP), subgrouped following the results from two clinical tests with high validity and reliability, differ in demographic characteristics and weekly amount of days with bothersome symptoms through the second half of pregnancy.

Design A prospective longitudinal cohort study.

Participants Pregnant women with pelvic and lumbopelvic pain due for their second-trimester routine ultrasound examination.

Setting Obstetric outpatient clinic at Stavanger University Hospital, Norway.

Methods Women reporting pelvic and lumbopelvic pain completed a questionnaire on demographic and clinical features. They were clinically examined following a test procedure recommended in the European guidelines for the diagnosis and treatment of PGP. Women without pain symptoms completed a questionnaire on demographic data. All women were followed weekly through an SMS-Track survey until delivery.

Primary and secondary outcome measures The outcome measures were the results from clinical diagnostic tests for PGP and the number of days per week with bothersome pelvic pain.

Results 503 women participated. 42% (212/503) reported pain in the lumbopelvic region and 39% (196/503) fulfilled the criteria for a probable PGP diagnosis. 27% (137/503) reported both the posterior pelvic pain provocation (P4) and the active straight leg raise (ASLR) tests positive at baseline in week 18, revealing 7.55 (95% CI 5.54 to 10.29) times higher mean number of days with bothersome pelvic pain compared with women with both tests negative. They presented the highest scores for workload, depressed mood, pain level, body mass index, Oswestry Disability Index and the number of previous pregnancies. Exercising regularly before and during pregnancy was more common in women with negative tests.

Conclusion If both P4 and ASLR tests were positive mid-pregnancy, a persistent bothersome pelvic pain of more than 5 days per week throughout the remainder of pregnancy could be predicted. Increased individual control over work situation and an active lifestyle, including regular exercise before and during pregnancy, may serve as a PGP prophylactic.

INTRODUCTION

Pelvic girdle pain (PGP) during pregnancy affects approximately half of all pregnant women, and for 25%–30% the condition becomes severe.1,2 The aetiology of PGP is still unknown, and the underlying mechanisms have not been fully investigated.1,2 Researchers have explored the physical, psychological and socioeconomic implications of PGP during pregnancy.3 Pain-related restrictions on physical activity have been described, both during pregnancy and after childbirth, and the psychological impact on perceived health, sexual life and quality of life has been explored, as well as the prevalence of sick leave due to PGP.3,6

PGP is classified into specific (caused by trauma) or non-specific (multifactorial).3 Several clinical tests are needed to diagnose the latter, including pain provocation and functional ability tests. However, there is still no ‘gold standard’ for diagnosing PGP. The European guidelines present evidence-based
recommendations for the diagnosis and treatment of PGP, but inconsistencies on the definition as well as treatment still prevail.3 7 Classification of PGP can, according to guidelines, only be reached after lumbar causes have been excluded through a clinical examination.7 All tests recommended in the European guidelines have a very high specificity, but generally a low sensitivity. Hence, it is recommended to perform all the tests, as one negative test is not sufficient to rule out PGP.7 The posterior pelvic pain provocation test (P4), for diagnosing sacroiliac joint dysfunction, and the active straight leg raise test (ASLR), for detecting failing force closure, have shown high validity and reliability.5–10

In a Swedish study, substantial agreement between examiners using ASLR and P4 tests was found in discriminating non-specific lumbopelvic pain into lumbar pain and PGP in pregnant women.11 Together with a description of pain location, these tests are considered relevant when evaluating affliction in pregnant women likely to have PGP.12

So far, the longitudinal course of PGP in pregnancy is incompletely examined. In prospective studies data are usually collected at baseline and at one or a few follow-ups. Measuring only at a few points in time may indicate stability in the examined condition, and a fluctuating course may be missed. A difference could reflect only a temporary fluctuation in an otherwise stable condition. Accordingly, a more frequent data collection is warranted to accurately describe the clinical course. Mobile phones and text messages have previously been found feasible when collecting frequent longitudinal data in clinical settings.13–15 Phones are usually at hand in daily life; hence, this method yields a high response rate for weekly measures.

The objective of this study was to explore if pregnant women with probable PGP, subgrouped following the results from two valid and reliable clinical tests recommended in the European guidelines, differ in demographic and clinical characteristics at mid-pregnancy and in weekly amount of days with bothersome symptoms through the second half of pregnancy. The hypothesis was that sacroiliac dysfunction and failing force closure diagnosed at mid-pregnancy may predict a course of bothersome symptoms through the second half of pregnancy.

METHODS

This is a prospective longitudinal cohort study of pregnant women who had their second-trimester routine ultrasound examination in pregnancy week 18 at an obstetric outpatient clinic at Stavanger University Hospital, Norway, from mid-March to mid-June 2010. At the hospital, all the women were asked by a midwife about their experience of pain in the lumbopelvic region. The inclusion criteria were current lumbopelvic pain or isolated pelvic pain, singleton pregnancy and good proficiency in the Norwegian language. Women who met the criteria were informed about the study, handed a letter of consent to fill in if they agreed to participate, and an envelope with questionnaires on demographic and clinical data to complete at home. An appointment with a chiropractor for a physical examination was arranged, and the women were asked to bring the completed questionnaires with them to the consultation. Women without pain symptoms were informed about the study, handed a letter of consent to fill in if they agreed to participate, and a questionnaire on demographic data to complete and leave at the reception on departure. All consenting women were followed from week 18 with weekly, automated text messages (SMS-Track).

Two chiropractors (SM and IK) performed a physical examination of the pelvic region, including diagnostic tests recommended in the European guidelines for the diagnosis and treatment of PGP, and a neurological examination of the lower extremities.7

Sequence of stability and pain provocation tests for PGP

Active straight leg raise

The test is performed with the patient in a supine position with a straight leg and the feet 20 cm apart. The test is performed after the instruction ‘try to raise your legs, one after the other, above the couch for 20 cm without bending the knee’. The patient is asked to score impairment on a 6-point scale: not difficult at all=0; minimally difficult=1; somewhat difficult; difficult=2; fairly difficult=3; very difficult=4; unable to do=5. The scores on both sides are added, so that the total score range from 0 to 10.8

Gaenslen’s test

The patient, lying supine, flexes the knee and hip of the same side, the thigh being crowded against the abdomen with the aid of both the patient’s hands clasped about the flexed knee. The patient is then brought well to the side of the table, and the opposite thigh is slowly hyper-extended by the examiner with gradually increasing force by pressure of the examiner’s hand on top of the knee. With the opposite hand, the examiner assists the patient in fixing the lumbar spine and pelvis by applying pressure over the patient’s clasped hands. The test is positive if the patient experiences pain, either local or referred on the provoked side.10

Long dorsal sacroiliac ligament test

The subject lies on her side with slight flexion in both the hip and knee joints. If the palpation causes pain that persists more than 5 s after removal of the examiner’s hand, it is recorded as pain. If the pain disappears within 5 s, it is recorded as tenderness.17

Modified Trendelenburg’s test

The patient stands on one leg, and flexes the other at 90° in the hip and knee. If pain is experienced in the symphysis, the test is positive.17

Patrick’s FABER test

The subject lies supine. One leg is flexed, abducted and externally rotated (FABER, abbreviation of flexion
abduction and external rotation) so that the heel rests on the opposite knee. If pain is felt in the sacroiliac joints or in the symphysis, the test is considered positive.17

**Posterior pelvic pain provocation test**

The test is performed with the woman supine and the hip flexed to an angle of 90° on the side to be examined: a light manual pressure is applied to the patient’s flexed knee along the longitudinal axis of the femur while the pelvis is stabilised by the examiner’s other hand resting on the patient’s contralateral superior anterior iliac spine. The test is positive when the patient feels a familiar well-localised pain deep in the gluteal area on the provoked side. A similar test is described as posterior shear or ‘thigh trust’.17,18

**Symphysis palpation test**

The subject lies supine. The entire front side of the pubic symphysis is palpated gently. If the palpation causes pain that persists more than 5 s after removal of the examiner’s hand, it is recorded as pain. If the pain disappears within 5 s, it is recorded as tenderness.17

A demographic questionnaire used in an earlier study on pelvic pain in pregnancy was filled in at baseline.19 The women marked the pain location on drawings with the pelvis and the low back demarcated. Pain intensity was rated on a Numerical Rating Scale (NRS) from 0 to 100, where 0 meant ‘No pain’ and 100 ‘Unbearable pain’. Information on pain-related activities of daily living (ADL) was collected through the Oswestry Disability Index (ODI), which at the time of the data collection (ADL) was collected through the Oswestry Disability Index (ODI), which at the time of the data collection was one of the principal outcome measures for defining disabling effects from spinal disorders and PGP.7 20 It is a patient-completed questionnaire which gives a subjective percentage score of the level of function (disability) in 10 ADLs in patients with low back pain. Every activity contains six statements on how it is performed. The statements are scored from 0 to 5, with the first statement scoring 0 through to the last at 5. The scores for all questions answered are summed, then multiplied by 2 to obtain the index (range 0–100). Zero is equated with no disability and 100 is the maximum disability possible.

Physical workload was measured through five answer categories ranging from ‘sedentary’ to ‘heavy’, following a scale used in Stockholm Public Health questionnaire.21 The question on job satisfaction was a bipolar 5-point Likert scale with increments in two opposite directions (‘Very bad’ and ‘Very good’) and a neutral point in the middle.22

Every Sunday the women were asked through a short message service (SMS) how many days the previous week they had experienced bothersome pelvic pain: ‘How many days during the previous week has your pelvic pain been bothersome, (ie, affected your daily activities or routines)?’ If there was no reply, the question was repeated 24 hours later. The question should be answered with one single number between 0 and 7. The response was automatically entered into a database, which collected the continuous information from each participant over the duration of the study.

Demographic descriptive data are presented as median values with IQRs for continuous variables, and as frequencies for categorical variables. For univariate comparisons between symptomatic and asymptomatic subgroups, the non-parametric Kruskal-Wallis statistics were used. Categorical predictors in our model were four groups following the outcome from the ASLR and P4 tests (1: P4 positive, 2: ASLR positive, 3: both P4 and ASLR positive, 4: ASLR and P4 negative), time (pregnancy week), and the interaction term between time and test group for investigating whether the trajectory of SMS-reported number of bothersome days differed between the test groups. Other predictors in the model were age, number of previous deliveries and body mass index (BMI) before pregnancy.

The longitudinal trajectory of the SMS-Track response was modelled using a generalised estimating equations (GEE) approach, extending the generalised linear model to correlated longitudinal data and clustered data within subjects. The within-subject dependencies resulting from repeated measurement were modelled assuming an autoregressive relationship in the working correlation matrix. As the outcome variable was count data (weekly number of bothersome days with pain), the Poisson distribution was assumed with a log-link function.

Data were analysed using SPSS V.22.0 software. The Strengthening the Reporting of Observational Studies in Epidemiology reporting guideline was used during the writing of this article.

**Patient and public involvement**

Patients and the public were not involved in developing the research questions, outcome measures, as well as in the design and conduct of the study, or in the recruitment of patients.

**RESULTS**

Overall, 506 women agreed to participate in this study. Three were excluded due to incomplete data. On ultrasound examination in pregnancy week 18 did 42% (212/503) of the women report pain in the lumbopelvic region. A clinical examination revealed that 39% (196/503) of the women fulfilled the criteria for a probable PGP diagnosis, and 27% (137/503) showed positive response to ASLR and P4 tests. A further 12 women reported pelvic pain but did not respond to recommended clinical tests, and were therefore placed in the ‘ASLR and P4 tests negative group’.

There were significant differences in some demographic and clinical features at baseline between the women with and without pelvic pain and with different test outcomes (tables 1–2).

Women with positive P4 and ASLR tests experienced heavier workload. They also presented higher BMI at week 18, exercised less both before and during pregnancy, and slightly more than one-third reported feeling depressed.
| Variables                              | No pelvic pain | P values a–b | P4 and ASLR tests negative | P values a–c | P values b–c | P4 only c | P values a–d | P values b–d | P values d | P values a–e | P values b–e | Both ASLR and P4 | P values a–f | P values b–f | P values e | All ASLR and P4 f |
|---------------------------------------|----------------|--------------|----------------------------|--------------|--------------|-----------|--------------|--------------|-----------|--------------|--------------|------------------|--------------|--------------|-------------|-----------------|
| % (n)                                 | 58 (291)       | 15 (75)      | 36 (50)                    | 30.5 (26.7/33.2) | 0.92         | 0.21       | 30.0 (26.0/33.0) | 0.30         | 0.10       | 28.0 (24.0/34.0) | 0.44         | 0.11         | 29.0 (25.7/33.0) | 0.46         | 0.36         | 15.0 (12.0/17.0) |
| Age, median (IQR)                     | 29.0 (26.0/34.0) | 0.28         | 30.0 (28.0/34.0)           | 0.44         | 0.60         |           | 30.5 (26.7/33.2) | 0.92         | 0.21       | 30.0 (26.0/33.0) | 0.30         | 0.10         | 28.0 (24.0/34.0) | 0.44         | 0.11         | 29.0 (25.7/33.0) |
| Education years, median (IQR)         | 15.0 (13.0/17.0) | 0.64         | 15.0 (13.0/17.0)           | 0.55         | 0.89         |           | 15.0 (12.0/17.0) | 0.55         | 0.49       | 15.0 (12.0/17.0) | 0.41         | 0.33         | 15.0 (12.0/16.2) | 0.46         | 0.36         | 15.0 (12.0/17.0) |
| Physical workload, median (IQR)       | 3.0 (1.0/4.0)  | 0.01         | 2.0 (1.0/4.0)              | 0.00         | 0.24         |           | 3.0 (2.0/4.0)    | 0.96         | 0.00       | 3.0 (2.0/3.0)    | 0.00         | 0.00         | 4.0 (3.0/4.0)    | 0.01         | 0.01         | 3.0 (2.0/4.0)    |
| Work satisfaction, median (IQR)       | 5.0 (4.0/5.0)  | 0.32         | 4.0 (4.0/5.0)              | 0.88         | 0.66         |           | 4.5 (4.0/5.0)    | 0.96         | 0.80       | 5.0 (4.0/5.0)    | 0.91         | 0.74         | 4.0 (4.0/5.0)    | 0.52         | 0.38         | 4.5 (4.0/5.0)    |
| BMI, median (IQR)                     | 23.9 (22.3/27.0) | 0.03         | 24.9 (22.7/27.9)           | 0.03         | 0.33         |           | 25.6 (23.7/29.9) | 0.02         | 0.44       | 26.0 (24.0/30.4) | 0.08         | 0.52         | 25.0 (22.0/28.0) | 0.01         | 0.47         | 25.6 (22.8/28.5) |
| Number of births, median (IQR)        | 1.0 (0.0/1.0)  | 0.15         | 1.0 (0.0/2.0)              | 0.10         | 0.65         |           | 1.0 (0.0/1.5)    | 0.05         | 0.39       | 1.0 (0.0/2.0)    | 0.15         | 0.43         | 1.0 (0.0/1.0)    | 0.07         | 0.98         | 1.0 (0.0/1.0)    |
| Exercising before pregnancy, % (n)    | 37 (108)       | 0.19         | 36 (27)                    | 0.21         | 0.86         |           | 34 (17)       | 0.69         | 0.60       | 37.5 (9)      | 0.11         | 0.42         | 27 (17)         | 0.07         | 0.45         | 31 (43)         |
| Exercising during pregnancy, % (n)    | 20 (60)        | 0.01         | 19 (14)                    | 0.03         | 0.16         |           | 8 (4)         | 0.47         | 0.37       | 16 (4)        | 0.07         | 0.22         | 11 (7)          | 0.00         | 0.12         | 11 (15)         |

IQR (25/75). P value from non-parametric Kruskal-Wallis statistics. Subgroups marked with lower case letters from a to f for identification of p value from non-parametric test. ASLR, active straight leg raise test; BMI, body mass index; P4, posterior pelvic pain provocation test.
| Variables                                      | ASLR and P4 tests positive | P4 and ASLR tests negative | P4 only | ASLR only | Both ASLR and P4 | All ASLR and P4 |
|-----------------------------------------------|-----------------------------|-----------------------------|---------|-----------|-----------------|-----------------|
| % (n)                                         |                             |                             |         |           |                 |                 |
| No pelvic pain a                              | 58 (291)                    | 15 (75)                     | 36 (50) | 18 (24)   | 46 (63)         | 27 (137)        |
| &nbsp;&nbsp;&nbsp;&nbsp;Sick leave, % (n)       |                             |                             |         |           |                 |                 |
| &nbsp;&nbsp;&nbsp;&nbsp;&nbsp;P4 and ASLR tests positive | 25 (74)                    | 30 (23)                     | 38 (19) | 56 (14)   | 79 (51)         | 84 (59)         |
| &nbsp;&nbsp;&nbsp;&nbsp;P4 and ASLR tests positive |                             |                             |         |           |                 |                 |
| &nbsp;&nbsp;&nbsp;&nbsp;P4 only | 20 (60)                    | 21 (16)                     | 30 (15) | 36 (9)    | 01 (01)         | 01 (01)         |
| &nbsp;&nbsp;&nbsp;&nbsp;ASLR only | 13 (37)                    | 45 (34)                     | 62 (21) | 54 (13)   | 0.53 (38)       | 0.50 (42)       |
| &nbsp;&nbsp;&nbsp;&nbsp;PP previous pregnancies, % (n) | 4 (12)                    | 24 (18)                     | 93 (12) | 99 (6)    | 0.97 (21)       | 0.87 (23)       |
| &nbsp;&nbsp;&nbsp;&nbsp;PP before pregnancy, % (n) |                             |                             |         |           |                 |                 |
| &nbsp;&nbsp;&nbsp;&nbsp;ODI week 18, median (IQR) | 16.0 (9.0/26.0)            | 0.51 (12.0/26.0)            | 0.00    | 22.0 (16.0/32.0) | 0.00 (28.0 (18.0/34.0) |                 |
| &nbsp;&nbsp;&nbsp;&nbsp;PP week 18, median (IQR) | 20.0 (0.0/40.0)            | 0.00 (20.0/62.0)            | 0.00    | 40.0 (27.7/57.5) | 0.00 (50.0 (25.0/75.0) |                 |
| &nbsp;&nbsp;&nbsp;&nbsp;ASLR score, median (IQR) | 16.0 (9.0/26.0)            | 0.51 (12.0/26.0)            | 0.00    | 22.0 (16.0/32.0) | 0.00 (28.0 (18.0/34.0) |                 |
| &nbsp;&nbsp;&nbsp;&nbsp;LDL test positive, % (n) | 33.3 (25)                   | 0.01 (38)                   | 0.04    | 75 (18)   | 0.01 (84)       | 0.00 (80)       |
| &nbsp;&nbsp;&nbsp;&nbsp;Gaënslen’s test positive, % (n) | 5.3 (4)                    | 0.01 (14)                   | 0.00    | 21 (5)    | 0.00 (52)       | 0.00 (38)       |
| &nbsp;&nbsp;&nbsp;&nbsp;Symphysis test positive, % (n) | 29.3 (22)                   | 0.54 (27)                   | 0.41    | 63 (15)   | 0.31 (60)       | 0.16 (58)       |
| &nbsp;&nbsp;&nbsp;&nbsp;Modified Trendelenburg’s test positive, % (n) | 6.7 (5)                    | 0.83 (8)                    | 0.40    | 17 (4)    | 0.23 (21)       | 0.29 (17)       |
| &nbsp;&nbsp;&nbsp;&nbsp;FABER test positive, % (n) | 4.0 (3)                    | 0.01 (13)                   | 0.00    | 33 (8)    | 0.00 (41)       | 0.00 (34)       |
| &nbsp;&nbsp;&nbsp;&nbsp;SMS-Track, median (IQR) | 3.0 (2.0/6.0)               | 0.00 (3.0/7.0)              | 0.00    | 4.5 (2.0/6.0) | 0.00 (6.0 (4.0/7.0) |                 |

IQR (25/75). P value from non-parametric Kruskal-Wallis statistics. Subgroups marked with lower case letters from a to f for identification of p value from non-parametric test. ASLR, active straight leg raise test; Faber, flexion abduction and external rotation; LDL, long dorsal ligament test; NA, not applicable; ODI, Oswestry Disability Index; P4, posterior pelvic pain provocation test; PP, pelvic pain.
during the pregnancy. Physical disability (ODI) and pain level (NRS) at week 18 were considerably higher in women with positive tests than in women reporting pain but having negative P4 and ASLR tests (table 2).

Women with a positive ASLR, but negative P4 test, had the highest number of previous pregnancies. Almost half of the women with both P4 and ASLR tests positive had been on sick leave during their pregnancy. Apart from the P4 and ASLR, the long dorsal ligament test showed the highest positive response rate, followed by the symphysis provocation test (table 2).

The SMS-Track response rate was 75% (2148 responses to 2877 sent messages). Due to a declining SMS-Track response at the end of the pregnancy, we stopped our SMS-Track analysis at week 38. A GEE analysis revealed that all entered variables, except ‘age’, were significant predictors for the number of days with bothersome pelvic pain, and there was a significant interaction between diagnostic group and time, implying that the time course of days with bothersome pelvic pain was different for the different test groups.

The estimated weekly mean number of days with bothersome pelvic pain for the different test groups is presented in figure 1. Women with both P4 and ASLR tests positive experienced from week 18 a high weekly mean number of days (≈5) with bothersome pelvic pain throughout the pregnancy. Women with both tests negative showed a steadily rising number of bothersome days throughout the pregnancy, from 0.5 day in week 18, to 2 days in week 37. The group with a P4 positive and an ASLR negative test had approximately 3 days of bothersome pelvic pain in week 18, which was considerably lower than the group with both tests positive, but showed rapidly increasing number of days with pain. In week 29, the number of days with bothersome pelvic pain equaled that of the group with both tests positive, and thereafter matched this group. Women with a positive ASLR and a negative P4 test also showed 3 days of bothersome pelvic pain in week 18, but never reached the mean number of bothersome days reported by women with P4 and both tests positive.

The parameter estimates output showed the estimated rate for experiencing bothersome days to be 7.5 times higher in women with both ASLR and P4 tests positive, compared with the estimate for women with both tests negative. Women with both tests positive were estimated with twice the amount of bothersome days per week (table 3). For women with either P4 or ASLR test positive and a lower incidence rate, the mean amount of bothersome days was lower, but still estimated as approximately 1.5 times higher than for women with both tests negative. For every pregnancy, the mean number of bothersome days increased by 13.5%. Even a slightly higher BMI had a significant impact on the mean amount of bothersome days. Age had no impact.

Figure 1  Estimated mean number of bothersome days in the latter half of pregnancy. If both P4 and ASLR tests were positive at week 18 (blue), the highest mean weekly number of days with bothersome pelvic pain throughout the second half could be expected. If both tests were negative, the mean number of bothersome days would never reach the amount reported by women with positive tests (purple). Women with a positive P4 test and a negative ASLR test (green) presented approximately 3 days of bothersome pelvic pain in week 18, with the mean number of days increasing rapidly until week 29, equaling the group with both tests positive (blue). Interestingly, women with a positive ASLR and a negative P4 test (red) also showed 3 days of bothersome pelvic pain at week 18, but never reached the mean number of bothersome days reported by women with P4 and both tests positive. ASLR, active straight leg raise test; P4, posterior pelvic pain provocation test.
DISCUSSION

To our knowledge, this is the only study in which women with pelvic pain in pregnancy have been followed with SMS-Track. The main result of this study was that if both P4 and ASLR tests were positive in pregnancy week 18, a persistent pelvic pain of more than 5 days/week throughout the remainder of pregnancy could be predicted. If either test was positive in week 18, a similar course was shown, but women with a positive P4 test revealed a more uncomfortable course than women with a positive ASLR test, who never reached the bothersome levels of the other groups. Robinson and coworkers reported a similar outcome for the P4 test in a prospective cohort study on the association between sociodemographics, psychological and clinical factors measured at mid-pregnancy, and disability and pain intensity at week 30. However, their data showed no significant association between the ASLR test result and the disability and pain intensity in pregnancy week 30.

Although women who had a positive ASLR test and negative P4 test at baseline presented a comparatively low mean number of bothersome days with pain, they also had the highest mean number of previous pregnancies and the highest mean rate of pelvic pain in previous pregnancies. Interestingly, our data also revealed that they exercised more frequently in comparison with women in the other positive test groups, both before and during the present pregnancy.

Interpretation

Since sufficient force closure of the sacroiliac joints requires appropriate muscular, ligamentous and fascial interaction, may women with pelvic pain in previous pregnancies have experienced that exercising improves muscle activation, recovers function and decreases pain. Additionally, experiences of pain prevention and rehabilitation in previous pregnancies may work as an incitement to engage in physical activity and regular exercise, both before and during pregnancy.

Our analysis also revealed a significant difference between the test groups in women described feeling depressed, and that a prepregnancy BMI slightly higher than average had a significant impact on the mean number of bothersome days. Distress has previously been identified as a factor associated with a higher likelihood of PGP in pregnancy, as have a higher BMI and a higher gestational age. One previous study found distress contributing to disability, but not to pain intensity.

Nevertheless, some individuals seem to tolerate pain better, have less catastrophising tendencies and show more positive social response to pain, regardless of exposure to stressful circumstances and/or internal distress. Finally, women with a possibility to control their own work situation have better health during pregnancy than women without such chances. As indicated in this study and confirmed in previous studies, most pregnant women benefit from exercise since it increases pain tolerance, improves or maintains physical fitness, helps with weight management, reduces the risk of gestational diabetes in obese women, and enhances psychological well-being.

Limitations

A limitation of this study is the retrospectively collected information on pain in previous pregnancies and pain before pregnancy, which may produce biased results. Another limitation was found in the data collection via the SMS-Track system. In Norway, at the time of the study, there were more mobile phone service providers than in neighbouring countries, where SMS-Track studies previously had been successfully performed. Unfortunately, we had frequent problems reaching women through some of the providers. These problems led to data missing at random, but the GEE analytic approach may in these instances compensate for missing data. However, using the SMS-Track system in data collection is also a strength in our study, since it provides instant data on participants’ situation, and responses are immediately recorded in a datasheet, which minimises further data handling and risk of error.

CONCLUSION

If both ASLR and P4 tests are positive at a clinical examination in mid-pregnancy, a course of persistent
bothersome pelvic pain for more than 5 days per week throughout pregnancy may be predicted. The number of days per week with bothersome pelvic pain increases for every added pregnancy, but individual control over work situation and regular exercise may work as a PGP prophylactic since it invigorates a positive impact on optimal force closure of the pelvis, reduces risk of instability in the pelvic joints and enhances overall well-being.

Since there is still no gold standard for diagnosing PGP, particularly regarding the number of tests at the clinical examination, we recommend further research in this area, aiming at predictive, preventive and diagnostic measures for identifying women at risk of developing PGP in pregnancy. It would, for example, be interesting to see if women with a history of PGP have a higher pain-related anxiety and if it influences pain.

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Contributors SM, IK, JPL, KA and IØ made substantial contributions to the conception and design of the work and revising it critically for important intellectual content. SM and IK performed the data collection. SM, IK and AMG controlled the data. SM and KSB analysed the data. SM, IK, KA, AMG, KSB and IØ made final approval of the version to be published. SM agrees to be accountable for all aspects of the work in ensuring that questions related to the accuracy or integrity of any part of the work are appropriately investigated and resolved.

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Competing interests None declared.

Patient consent Obtained.

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