Statin treatment increases the clinical risk of tendinopathy through matrix metalloproteinase release – a cohort study design combined with an experimental study

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Recent experimental evidence indicates potential adverse effects of statin treatment on tendons but previous clinical studies are few and inconclusive. The aims of our study were, first, to determine whether statin use in a cohort design is associated with tendinopathy disorders, and second, to experimentally understand the pathogenesis of statin induced tendinopathy. We studied association between statin use and different tendon injuries in two population-based Swedish cohorts by time-dependent Cox regression analysis. Additionally, we tested simvastatin in a 3D cell culture model with human tenocytes. Compared with never-users, current users of statins had a higher incidence of trigger finger with adjusted hazard ratios (aHRs) of 1.50 for men (95% confidence interval [CI] 1.21–1.85) and 1.21 (1.02–1.43) for women. We also found a higher incidence of shoulder tendinopathy in both men (aHR 1.43; 1.24–1.65) and women (aHR 1.41; 0.97–2.05). Former users did not confer a higher risk of tendinopathies.

In vitro experiments revealed an increased release of matrix metalloproteinase (MMP)-1 and MMP-13 and a weaker, disrupted matrix after simvastatin exposure. Current statin use seems to increase the risk of trigger finger and shoulder tendinopathy, possibly through increased MMP release, and subsequently, a weakened tendon matrix which will be more prone to injuries.

Statins are commonly used to lower blood cholesterol levels, which helps prevent cardiovascular disease1–3. Although they are generally believed to be safe, muscle toxicity is not uncommon4,5. Increasing attention has also been directed towards potential harmful effects on tendons, leading to tendinopathy or rupture. Case reports and pharmacovigilance data suggest an increased risk of tendon rupture and tendinopathy in statin users4–8, especially in men7,9; however, prospective cohort studies on the adverse effects of statin therapy are scarce and inconclusive10–14. The initial pharmacovigilance study describing a relationship between statins and tendon impairment reported “tendinitis” as the most frequent tendon adverse event with the highest incidence during the first year of statin use7. Simvastatin, is the most prescribed statin in Sweden and has together with atorvastatin and lovastatin the highest risk for muscle related side-effects, due to nonselective abilities to diffuse into other tissues than the liver5.

Tendinopathy, also known as tendinosis or tendinitis, is a broad term used to describe tendon degeneration15. The condition is characterized by collagen degeneration, fiber disorganization and increased amounts of ground substance. We have previously shown that trigger finger tendons show a histological appearance and gene expression pattern similar to tendinopathy at other locations, suggesting that this condition might be the most common...
form of tendinopathy. Because it is clinically an easily defined tendinopathy, it can serve as a suitable outcome variable in an observational study.

An exact molecular mechanism of action for statins in tendon cells has not been established. In vitro studies have shown that extracellular matrix strength is reduced after statin treatment but, surprisingly, without altering the total levels of collagen. This finding indicates that alterations in the balance of matrix metalloproteinases (MMPs) might play a role.

Given the indeterminate evidence on statin use and the risk of tendon pathology, we used two large Swedish population-based cohorts to evaluate a potential association between statin use and the risk of tendinopathy. We chose to study whether statins are associated with a higher risk of trigger finger and with tendinopathy in the shoulder or the Achilles tendon. Moreover, we used an in vitro model with artificial tendons (made from human tendon fibroblasts) to study the possible role of statin-driven MMP release in association with a weakened extracellular matrix.

**Results**

To study the involvement of statin use in the development of tendon disorders a cohort study was performed with a time-dependent Cox regression analysis using two Swedish population-based cohorts in combination with three national registers. The baseline characteristics of the study participants are shown in Table 1. The mean age at baseline was slightly <70 years. Statin use was common in both women (37%, 19,323) and men (44%, 17,854). The most frequent statin prescribed was simvastatin (69%), followed by atorvastatin (24%), rosuvastatin (4%) and pravastatin (2%).

|                       | SMC (n = 52,220 women) | COSM (n = 40,713 men) |
|-----------------------|------------------------|-----------------------|
| Number of women/men   | 19,323                 | 17,854                |
| Age, mean (SD)        | 68.4 (7.8)             | 69.5 (8.5)            |
| Body mass index, mean (SD) | 25.6 (4.0)             | 24.7 (3.9)            |
| Metabolic equivalents (METs, in 1998) | 42.4 (4.8)             | 42.5 (4.7)            |
| Charlson weighted comorbidity index, mean (SD) | 0.29 (0.72)             | 0.23 (0.64)            |
| Energy intake (kcal/day) | 1715 (577)             | 1731 (582)            |
| Pack-years of smoking, mean (SD) | 11.5 (10.5)             | 10.6 (10.1)            |
| Education, n (%)      |                        |                       |
| <9 years              | 12,751 (70.0)          | 10,078 (72.0)         |
| 10–12 years           | 3175 (16.4)            | 1815 (13.0)           |
| >12 years             | 2585 (13.4)            | 2045 (14.6)           |
| Vocational education  | 38 (0.2)               | 57 (0.4)              |
| Corticosteroid use, n (%) | 1010 (5.2)            | 3538 (19.8)           |
| Quinolone use, n (%)  | 2962 (15.3)            | 6062 (34.0)           |
| Diabetes mellitus, n (%) | 2618 (13.6)           | 3963 (22.2)           |
| Renal insufficiency, n (%) | 455 (2.4)              | 1002 (5.6)            |
| Statin use, n (%)     |                        |                       |
| Simvastatin           | 13,266 (68.7)          | 12,256 (68.7)         |
| Pravastatin           | 415 (2.2)              | 243 (1.4)             |
| Fluvastatin           | 43 (0.2)               | 26 (0.2)              |
| Atorvastatin          | 4325 (22.4)            | 4584 (25.7)           |
| Rosuvastatin          | 775 (4.0)              | 745 (4.2)             |

Table 1. Descriptive characteristics of statin users and never-users. *% in statin user.

Statin treatment is associated with an increased risk of trigger finger. During 833,390 person-years of follow-up, we identified 1056 incident cases of trigger finger according to our ICD code criteria (626 women and 430 men). The incidence rates for trigger finger were 1.4/1000 person-years in women and 1.5/1000 in men. Compared with non-users, current users of statins conferred an overall higher risk of trigger finger: multivariable-adjusted hazard ratio (aHR) 1.50 (95% CI 1.21–1.85) for men and 1.21 (95% CI 1.02–1.43) for women (Fig. 1 and Table 2). The higher risk did not remain in former users. We found no relation between the duration of current statin use and the risk to develop trigger finger, i.e., the aHRs were similar in patients with 0–1 years versus >3 years of statin use. The highest apparent relative risk was seen in male users of rosuvastatin (aHR 2.19; 95% CI: 1.33–3.62) though this estimate is based on few cases. We found no statistically significant heterogeneity between the estimates for the different statins (p = 0.44 in women and p = 0.36 in men) or between prescribed doses.

Ongoing statin use is also associated with an increased risk of shoulder tendinopathy. In the same cohorts we identified 1102 women and 957 men with shoulder tendinopathy (Fig. 2 and Table 3). We
observed a higher incidence of shoulder tendinopathy in men and women who were current users of statins (aHRs 1.43; 95% CI: 1.24–1.65 and 1.41; 95% CI 0.97–2.05, respectively) compared with men and women who were never-users. Former users of statins were not affected by a higher incidence of shoulder tendinopathy. Again, we found no statistically significant heterogeneity between the estimates for the different statins (p = 0.78 in women and p = 0.10 in men).

Statin use and achilles tendon pathology. In these two cohorts few of the participants had had Achilles tendinopathy or Achilles tendon rupture as a diagnosis. Tendonopathy was seen in only 88 women and 131 men and ruptures in only 71 women and 237 men, which limit the precision of our estimates. Still, there was a tendency of an increased risk for tendinopathy in men and women who had ever used statins. The aHR for men was 1.36 (95% CI: 0.92–2.00) and 1.25 (95% CI: 0.81–1.92) for women (Fig. 3). The corresponding aHR for an Achilles tendon rupture was 1.16 in men (95% CI: 0.88–1.54) and 1.41 in women (95% CI: 0.88–2.27). No further subanalyses were done because of the small number of observations.

Table 2. Hazard ratios (HR) and 95% confidence intervals (CIs) of trigger finger associated with statin use in women and men. aMultivariable-adjusted model adjusted for age, BMI, METs, Charlson index, pack-years of smoking, energy intake, oral corticosteroid use, quinolone antibiotic use, diabetes mellitus, renal insufficiency and education level.
Simvastatin in vitro triggers a release of MMP-1 and MMP-13. We used a three-dimensional (3D) cell culture model with artificial tendons, to mechanismically investigate whether MMPs were involved in the adverse effect of statins (Fig. 4A,B). Simvastatin administration for 7 days led to a reduction in maximum force and stiffness by approximately half (both p-values < 0.005) without altering the cross-sectional area (p = 0.28, Fig. 5). The material properties, peak stress and elastic modulus were also reduced (both p-values = 0.03). Protein analyses of the cell culture supernatant showed no overall increase in protein levels but specific increase in levels of MMP-1 by a 6-fold and MMP-13 by 1.3-fold after statin treatment (both p-values < 0.03), whereas levels of MMP-3 were virtually unchanged (Fig. 6). Histological images with hematoxylin and eosin (H&E) staining confirmed a more disrupted matrix appearance after simvastatin exposure (Fig. 4C–1).
tendons degrading collagen type I in tendons23, and have previously been suggested to play a role in the degradation of collagen in tendons.

The protein content in the supernatant was unaffected by simvastatin exposure. MMP-1 and MMP-13 are collagenases and have been shown to degrade collagen in tendons18. Our study confirmed increased protein levels of MMP-1 and MMP-13 in the cell culture supernatant, whereas MMP-3 protein expression remained unaltered. Additionally, the total protein content in the supernatant was unaffected by simvastatin but increased gene expression levels for MMP-1 and MMP-13 were observed.

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A clear, independent higher rate in the development of tendinopathy during statin use, including trigger finger, has not previously been described. However, a possible association between statin use and tendon complications has frequently been discussed5,9,10,13,19. Although tendinopathy (e.g., tendinitis) has been proposed as a possible common side effect of statins than ruptures7, previous cohort studies have concerned only a link between statin use and tendon ruptures and observed no overall effect of statins5,11–14.

Hypercholesterolemia is a potential risk factor for tendinopathy12,20–22, and therefore it can be hard to separate the effects of hypercholesterolemia from those of statins. Nevertheless, we found an increased risk of tendinopathy only in current statin users and not former users, which supports a role of statin use in tendinopathy development. It has previously been described that most patients experience their tendon problems within the first year after statin initiation5,12, with a median time to onset of 8 months after initiation7. We believe that this finding can explain the lack of duration effect in our study. The finding also indicates that tendinopathy might be triggered in predisposed individuals, similar to what has been suggested for muscle related side-effects1. A previous study based on pharmacovigilance data reported a median of 23 days for tendon problem manifestation to disappear after statin cessation7 and a re-occurrence after statin treatment was re-introduced7,8.

We found a higher risk of trigger finger and shoulder tendinopathy in current statin users but not in former users in two large population-based cohorts. Moreover, our in vitro studies confirmed an adverse effect of simvastatin on tendon extracellular matrix, as well as an increased release of the collagenases MMP-1 and MMP-13 by human tendon fibroblasts.

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We noted no association with a more pronounced risk after a longer treatment duration and no clear dose-response relationship. Our findings support case reports and pharmacovigilance data which have shown that 59% of the patients experienced problems within the first year of statin treatment7,9. A possible explanation for the weakened relation over time might be pre-disposition in certain patients to tendon-related side effects. Once these patients have been diagnosed in a closed cohort, fewer predisposed patients are at risk to develop symptoms during an expanded follow-up time. Pre-disposing factors, such as genetic susceptibility, need to be identified, as for muscle related problems1.

We were able to confirm a detrimental effect of simvastatin, the most commonly used statin, on artificial tendons from human tendon fibroblasts. Simvastatin is a lipophilic drug which can passively diffuse through the cell membrane into other tissues e.g. muscle and tendons3. Both structural and material properties of the tendons were reduced but the cross-sectional area was unaltered. A previous study using the same in vitro model showed no effect on collagen content after simvastatin administration but increased gene expression levels for MMP-1, MMP-3 and MMP-1314. Our study confirmed increased protein levels of MMP-1 and MMP-13 in the cell culture supernatant, whereas MMP-3 protein expression remained unaltered. Additionally, the total protein content in the supernatant was unaffected by simvastatin exposure. MMP-1 and MMP-13 are collagenases and the main enzymes degrading collagen type I in tendons23, and have previously been suggested to play a role in shoulder tendinopathy24–26. Although the levels of total collagen do not appear to be altered by simvastatin14, the appearance of the collagen could still have been affected. Our histological analysis confirmed this by showing a more disrupted appearance in the matrix after simvastatin exposure. Our results suggest that the weakened mechanical properties of tendon tissue could be caused by a disturbed matrix metalloproteinase balance. Furthermore, a weakened tendon tissue is likely more prone to injuries.

Our study design has several strengths. We used two population-based cohorts and register information for men and women rendering similar covariate information retrieved before the diagnosis of tendinopathies. Statin use and tendinopathies were identically defined in both cohorts, with complete identification of statin use and specialist physician first diagnosed cases by use of national registers (The Swedish National Patient register, the Swedish Prescribed Drug register and the Cause of death register) and the unique personal identification...
number provided to all Swedish residents. The Swedish Patient register has high accuracy with a positive predictive value of 97% for musculoskeletal diagnoses. Moreover, we used a time to event analysis that included a time-dependent exposure analysis limiting the likelihood of a reverse causation phenomenon and immortal time bias, a possible explanation for findings of apparent associations with different diseases by statins observed when using a cruder analytical approach.

Our study also has some limitations. We used data from the Swedish National Patient Register, a register that only includes diagnoses ascertained by specialized physicians and not by primary health care. Therefore, a diagnosis made only by a general practitioner will be overlooked and the number of actual cases is most likely underestimated. However, this possibility is unlikely to be related to statin use. Diagnostic errors are in a similar manner probably to be found in both users and non-users. Of note, we found the most evident association between statin use and trigger finger, which has a more characteristic clinical picture than the other tendinopathies investigated. This diagnosis is therefore probably less susceptible to misclassification. Additionally, due to low frequency of users, we had low to moderate precision for the estimation of associations for pravastatin.
fluvastatin and rosuvastatin. Confounding by indication is another concern, where it is plausible that patients who were prescribed statins had a different risk profile than those who did not receive statins. Using our study cohort data, we were able to account for several important potential covariates known to influence tendon tissues. However, we lack information about lipid profiles in these patients and hypercholesterolemia is a possible risk factor for tendon injuries.

In conclusion, statin use appears to be associated with a higher risk of developing tendinopathy, in both men and women. We suspect that this adverse effect might be linked to an excessive MMP release followed by a weakened tendon matrix.

Methods

Study design. Two independent cohorts, combined with three national registers, were used to evaluate if statin use is involved in the development of tendon disorders. This was combined with an experimental study on human tendon fibroblasts, in vitro.

Observational studies. The observational part is based on 92,933 participants from two population-based cohorts in Sweden, which have previously been described31,32: the Swedish Mammography Cohort (SMC) and the Cohort of Swedish Men (COSM) – both parts of the national research infrastructure SIMPLER.
(www.simpler4health.se). The study has been approved by the regional research ethical review board at the Karolinska Institutet, Stockholm (2009/1682-31; 2009/1935-32; 2013/1605-32/5; 2016/1924-32).

SMC. All women in Västmanland and Uppland Counties in Sweden, born between 1914 and 1948, were invited in 1987 through 1990 to participate in the SMC32. Of the 90 303 women invited, 74% (n = 66 651) accepted and completed a first self-administered questionnaire on diet, alcohol consumption, education, living conditions, body weight and height. In 1997, the 56 030 women who were still alive and living in the study area received a second, expanded self-administered questionnaire. This expanded version included information on smoking status, physical activity and other lifestyle factors. In this second phase 70% (n = 39 227) of the women responded.

COSM. All men in Västmanland and Örebro Counties in Sweden, born between 1918 and 1952, were invited in 1997 to participate in the COSM32. Of the 100 303 eligible men, 49% (n = 48 850) accepted and completed a self-administered questionnaire on diet, alcohol consumption, education, living conditions, body weight and height, physical activity, smoking habits and other lifestyle factors.

**Exposure definition.** Data on exposure to statins were retrieved from The Swedish Prescribed Drug Register. This register includes all prescriptions dispensed in Sweden since July 2005. All drugs in the register are classified using the Anatomical Therapeutic Chemical (ATC) system and the register is almost complete for the entire population in Sweden (patient identity data are missing for <0.3% of all items).

We defined statin exposure as any use of statins registered with ATC codes C10AA01 to C10AA08 from 1 July 1 2005 until 31 December 2014. Current users were analyzed in duration categories (<1, 1–2, 2–3, >3 years). If an individual did not collect a new prescription at the pharmacy 90 days after the previous doses would have been consumed, we regarded this individual as a former user from that point onwards. The total duration of use was calculated as the difference between the first day of dispensing and the last day of dispensing plus 3 months, excluding gaps in treatment. The drugs are normally dispensed every third month. The individual statins that were analyzed were pravastatin and rosuvastatin (both hydrophilic) and simvastatin, fluvastatin and atorvastatin (all lipophilic statins). We additionally classified statin dose per day into three categories: low dose (simvastatin 5–10 mg, pravastatin 10 mg, atorvastatin 10 mg, rosuvastatin 5–10 mg), medium dose (simvastatin 20–40 mg, pravastatin 20–40 mg, fluvastatin 20–40 mg, atorvastatin 20–40 mg, rosuvastatin 20 mg) and high dose (simvastatin 80 mg, fluvastatin 80 mg, atorvastatin 80 mg, rosuvastatin 40 mg).

**Outcome definition.** We defined the outcome as the first occurrence of any of our diagnosis in inpatient or outpatient specialist care data after the index date 1 July 2005 (starting the first day of the prescription register) until 31 December 2014. First occurrence of trigger finger, tendinopathy or Achilles tendon rupture was considered an outcome based on ICD-10 codes: M65.3 for trigger finger, M75.1, M75.2, M75.3, M75.4 and M75.5 for shoulder tendinopathy, M76.6, including sub-diagnosis under that coding for Achilles tendinopathy and S86.0, M66.3 H and M66.2 H for Achilles tendon rupture. Those patients with a tendinopathy diagnosed before the index date were excluded from the analysis.

**In vitro studies with tendon constructs.** We isolated tendon fibroblast by collagenase digestion from small segments of human semitendinosus tendons in five patients undergoing anterior cruciate ligament reconstructive surgery as previously described81. The experiments were approved by the regional ethics review board in Linköping, Sweden (2015/408-31) and patients gave written informed consent to participation. The investigation has been conducted according to the Declaration of Helsinki. Cells were seeded in flasks and cultured to confluence in DMEM/F12 supplemented with 10% fetal bovine serum (FBS) and 1% Penicillin-Streptomycin.

**Tendon construct formation.** Tendon constructs were assembled as previously described with small modifications81. The bottom of six-well plates were coated with silicon (SYLGARD, Dow-Chemicals) and two silk sutures (0.5 cm, Ethicon) were pinned onto this as anchor points (15 mm apart). Then, 250 000 cells (passage 2–5) were mixed into a fibrin gel and quickly spread in each well. The fibrin was left to set before it was covered with DMEM/F12 (supplemented with 10% FBS, 0.2 mM L-asorbic acid 2-phosphate, 0.05 mM L-Proline and 1% Penicillin-Streptomycin). The cell culture supernatant was replaced every second to third day and adhesions with DMEM/F12 supplemented with 10% FBS, 0.2 mM L-asorbic acid 2-phosphate, 0.05 mM L-Proline and 1% Penicillin-Streptomycin. The testing was performed in quadruplicate for each cell donor treated with simvastatin and in duplicate for each cell donor treated with DMSO.

**Mechanical testing of the constructs.** Tensile testing was performed in an Instron 3343 Single Column Testing System (10 N load cell) with a BioPuls2D® liquid chamber and Bluehill® material testing software (version 2.35.917, Instron, Buckinghamshire, United Kingdom). Construct diameter and length were measured before testing by capturing images with a digital camera at 11 cm from the construct. The diameter was measured in four places (the thickest and thinnest places on the right and left side of the construct) using ImageJ (NIH, USA). An average cross-sectional area was calculated assuming a circular cross section. Each picture also included a reference with known diameter and length. The sutures, in each end of the construct, were glued between sandpaper sheets to ensure adhesion within the clamp. The specimen was then transferred to a bath containing PBS at 30°C, and after a short adaptation period, the test was started. The samples were stretched at 4 mm/min until failure. The testing was performed in quadruplicate for each cell donor treated with simvastatin and in duplicate for each cell donor treated with DMSO.
MMP ELISA and total protein content measurement. Cell culture supernatant was stored at –70 °C until analysis. Levels of the collagenases (MMP-1 and MMP-13) and stromelysin 1 (MMP-3) were determined using sandwich ELISA kits (Catalogue #DM100, DM1300, DM300; R&D Systems) according to the manufacturer’s instructions. Total protein content in the supernatant was measured by the Bio-Rad protein assay according to the manufacturer’s instructions (Bio-Rad).

Histology. Constructs were fixed in 4% paraformaldehyde, followed by dehydration in a series of increasing concentrations of ethanol, before they were embedded in paraffin. The paraffin blocks were then sectioned longitudinally, in 7 μm sections and stained with routine haematoxylin and eosin. The sections were observed in a conventional light microscope.

Statistical analysis. Observational study. All cohort participants who were alive and resident in Sweden at the starting point (1 July 2005) were included in the analysis. We estimated age- and multivariable-adjusted hazard ratios (HRs) for relative risk of trigger finger, shoulder tendinopathy, Achilles tendinopathy and Achilles rupture in users versus non-users of statins using Cox proportional hazards regression analysis. We used years of observation as a time scale and with time-updated information on exposures. All individuals were followed from 1 July 2005 until date of outcome, date of death, emigration, or to the end of the observation period (i.e. 31 December 2014), whichever occurred first.

To minimize potential confounding a multivariable model – identified by a directed acyclic graph approach – was used that included the following variables: age (continuous), body mass index (weight [kg] divided by the height [m] squared, continuous), validated total physical activity (continuous as metabolic equivalent-hours/day) and educational level (<9 years, 10–12 years, >12 years and other education such as vocational) as a marker of socioeconomic status, smoking status categorized as never, former or current smoking, pack-years of smoking (continuous), total energy intake as an additional proxy variable for physical activity (continuous), weighted Charlson comorbidity index (continuous), corticosteroid use (ever/never), quinolone use (ever/never), diabetes mellitus (yes/no by self-report or by ICD-10 codes E10-E14 identified from the National Patient Register) and renal insufficiency (yes/no by ICD-10 codes N17-N19 identified from the National Patient Register). Analyses were performed with SAS 9.4. Lifestyle information was collected from the latest questionnaire cycle (i.e. 1997).

In vitro study. Results from mechanical testing, ELISA and protein content were analyzed with paired Student’s t tests (two-sided). The significance level was set at p < 0.05.

Data availability. Data cannot be made freely available as they are subject to secrecy in accordance with the Swedish Public Access to Information and Secrecy Act, but can be made available to researchers upon request (after subject to a review of secrecy and ethical approval) by contact to the national research infrastructure SIMPLER (Swedish Infrastructure for Medical Population-based Life-course and Environmental Research): www.simpler4health.se.

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Author contributions
P.E., P.A. and K.M. designed research. P.E., F.D., A.-C.L. and K.M. performed research. A.W. provided the cohorts used in this study and critiqued the manuscript. P.E., F.D., A.-C.L., P.A. and K.M. analyzed data and wrote the paper.

Competing interests
The authors declare no competing interests.

Additional information
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