Associations of Filaggrin Gene Loss-of-Function Variants and Human Papillomavirus-Related Cancer and Pre-Cancer in Danish Adults

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Abstract

**Purpose:** Filaggrin proteins are expressed in the skin, oral cavity, oesophagus, and cervical mucose. Loss-of-function mutations in the filaggrin gene (FLG) reduce filaggrin expression and cause an impaired skin barrier function. We hypothesized that FLG mutation carriers would be more susceptible to human papillomavirus (HPV) infection and thus a higher risk of HPV-related cancer and pre-cancer. We investigated the association of the FLG genotype with incidence of HPV-related cancer of cervix, vagina, vulva, penis, anus and head and neck, and pre-cancer of the cervix.

**Methods:** We included 13,376 persons from four population-based studies conducted in the same background population in Copenhagen, Denmark. Participants were genotyped for the most common FLG mutations in Europeans. Information on cancer was obtained from The Danish Cancer Registry until 11 July 2011.

**Results:** There were 489 cases of prevalent and 97 cases of incident HPV-related cancer and pre-cancer (median follow-up 11.5 years). There was a statistically significant association between FLG genotype and incident HPV-related cancer and precancer with a hazard ratio, HR = 2.1 (95% confidence intervals, CI: 1.2, 3.7) for FLG mutation carriers vs. wild types.

**Conclusions:** FLG loss-of-function mutations were associated with higher incidence of HPV-related cancers and pre-cancers that are potentially screening and vaccine preventable.

Introduction

Human papillomavirus (HPV) is a DNA virus infecting keratinocytes or cells in mucous membranes. Most known HPV types are largely harmless, some cause warts, whereas oncogenic types can cause pre-cancer and cancer of the cervix, vulva, vagina, penis, anus and a subgroup of head and neck cancers [1]. Precancerous lesions such as dysplasia and carcinoma in situ (CIS) of the cervix are common among women and if left untreated may lead to cancer. Cervical cancer is the fourth highest contributor to women cancer mortality worldwide and the second most common cause of cancer mortality among women in Africa [2]. Anal cancer primarily affects gay and bisexual men and is rare but increasing [1,3].

With the introduction of organized cervical cytological screening programs, cervical cancer incidence has been substantially reduced [4]. The developments in prophylactic HPV vaccination have renewed the interest in HPV-related cancers and cell changes. Vaccination against the two most important oncogenic HPV types (type 16 and 18) lowers the risk of anal, vulva, vaginal and penile infections with the two HPV types and decreases the risk of precancerous cervical lesions [5,6]. These two HPV types are likely responsible for 70% of cervical cancers and most of the non-cervical HPV-related cancers [1].

The epidermal layer of the skin provides a barrier against environmental exposures including microorganisms. Filaggrin proteins display structural and physiological functions in the skin but is also expressed in the oral cavity, cervix, endometrium, and vagina [7,8]. While the role of filaggrin outside the skin is largely unknown, the degradation products keep the epidermis acidic thereby preventing the colonization of microorganisms. Loss-of-function mutations in the filaggrin gene (FLG) reduce epidermal filaggrin levels and are among the most frequent known single-gene defects [9,10]. FLG loss-of-function mutations are strong genetic risk factors for atopic dermatitis in particular, but also...
Figure 1. Flowchart
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FLG loss-of-function mutations may lead to a greater susceptibility to HPV-related cancer and pre-cancer due to an impaired barrier function and atopic dermatitis [7,11]; elevated pH of the stratum corneum [7]; and a low grade skin inflammation [12–14]. We investigated the association of the FLG genotype and HPV-related cancer of cervix, vagina, vulva, penis, anus and head and neck and pre-cancer of the cervix according to the International Classification of Diseases (ICD) in four population-based studies.

**Material and Methods**

**Ethics statement**

Participants gave their informed written consent, and the studies were approved by the Ethics Committee of Copenhagen and the Danish Data Protection Agency. The recommendations of the Declaration of Helsinki were followed.

**Study populations**

We included the four population based studies, Monica10, Inter99, Health2006, and Allergy98, where the former three are recruited from the Danish Central Personal Register as random samples of the population in the southern part of the former Copenhagen County. The studies included questionnaires, physical examinations, and blood tests.

The Monica10 study was conducted in 1993-94 and included 2,656 persons of Danish origin (4,130 invited) between 40–71 years and had a participation rate of 64.3% [15].

The Inter99 study conducted in 1999-2001 included 6,784 persons aged 30–60 years [16]. The Inter99 study was a population-based randomized controlled trial (CT00289237, ClinicalTrials.gov) investigating the effects of lifestyle intervention on cardiovascular disease. The baseline participation rate was 52.5%. Details on the study and the intervention program have been described elsewhere [16]. Only participants with a Northern European origin were included in the current study. Both current and potential former nationalities of participants and their parents were considered (information from registries and self-reported questionnaires). A Northern European origin was defined as a Danish, Norwegian, Swedish, Icelandic, or Faroese nationality.

In the Health2006 study, a sample of 7,931 Danish citizens aged 18 to 69 years, born in Denmark, was invited to a general health examination [17]. A total of 3,471 (43.8%) individuals were examined between June 2006 and June 2008.

The Copenhagen Allergy study began in 1990 and included a group of persons randomly selected from the general population and a selected group of persons with allergic respiratory symptoms (recruited from a random sample of the general population by a screening questionnaire). We used data from the follow-up study in 1997–1998 (Allergy98) where a total of 1,966 persons aged 15–77 years with Danish nationality were invited for a health examination. A total of 1,216 (61.9%) participated [18].

We included a total of 13,376 persons with FLG genotype: 2,577, 6,247, 1,206, and 3,346 participants from the Monica10, the Inter99, the Allergy98, and the Health2006 study, respectively (Figure 1).

**FLG genotyping**

Individual regions covering the two most common null mutations of the FLG, R301X and 2282del4 (in the Inter99 and Health2006 studies also a region covering the R2447X mutation) were amplified from genomic DNA by allele-specific and asymmetric PCR using DNA tagged primers in all four studies.

The obtained PCR products were hybridized to MagPlex C microbeads (Lumixen, Austin, Texas) carrying the same tags as DNA probes [19]. Microbeads were subsequently analyzed on a Bio-Plex 200 (Bio-Rad Laboratories, Hercules, Calif). The filaggrin mutation analysis is ISO 15189 accredited. Samples also available as DNA were genotyped for FLG mutations. The genotyping success rates were: Monica10: 99.96%, Allergy98: ≥ 99%, Inter99: 99.95%, Health2006: 99.40%.

The gene frequencies were: Monica10: (R501X: 2.8%, 2282del4: 4.8%), Allergy98: (R501X: 3.4%, 2282del4: 3.8%), Inter99: (R501X: 3.4%, 2282del4: 4.4%, R2447X: 1.0%), and Health2006: (R501X: 3.4%, 2282del4: 4.7%, R2447X: 0.9%). All three FLG genotypes were in Hardy-Weinberg equilibrium (tested by the Hardy-Weinberg equilibrium test) [20]. FLG genotype was categorized as: no FLG mutations; or at least one FLG mutation (heterozygotes, homozygotes and compound homozygotes).

**Registry-based diagnoses**

People living in Denmark have since 1968 been assigned a unique and permanent personal civil registration number which enables linkage of data from complete national registers on an individual level. Information on cancer diagnoses was obtained from the Danish Cancer Register [21,22] according to the International Classification of Diseases (ICD). Reporting to the Cancer Registry has been mandatory since 1987. From 1943 to 1978 the Registry was classified according to the modified ICD-7, and from 1978 and onwards the diagnoses were coded in accordance with the ICD-10 [21]. Information on death from any cause and emigration status was obtained from the Danish Civil Registration System [23]. Participants were followed until 11 July 2011.

We included the cancers most strongly associated with HPV [24]: Cancer, cervix (ICD-7: 1711, 1712, ICD-10: C53), CIS, cervix (ICD-7: 5710, 5711, ICD-10: D06), dysplasia, cervix (ICD-7: 5712, ICD-10: N67), cancer, head and neck, subset (ICD-7: 1410, 1411, 1418, 1450, 1442, 1458, 1480, ICD-10: C01.9, C02.4, C02.8, C09.0, C09.1, C09.8, C09.9, C10.2, C10.8, C10.9, C14.0, C14.2, C14.8), cancer, vulva and vagina (ICD-7: 1760, 1761, 1762, 1763, 1765, ICD-10: C51, C32), cancer, penis (ICD-7: 1790, ICD-10: C60), and cancer, anus (ICD-7: No sufficiently specific ICD-7 code, ICD-10: C21.0–C21.8).

We defined three end points: cervical cancer and pre-cancer (Cervical dysplasia, CIS and cancer); all HPV-related cancers (subsets of cancers of head and neck, cervix, vulva, vagina, penis and anus as defined above); and all HPV-related cancers and pre-cancers (subsets of cancers of head and neck, cervix, vulva, vagina, penis and anus and cervical dysplasia and CIS).

Prevalent cancer and pre-cancer were defined as a diagnosis of the cancer and pre-cancer of interest before baseline, i.e. the date of the health examination (both ICD-7 and ICD-10 codes). Incident cancer and pre-cancer were defined as a diagnosis of the cancer and pre-cancer of interest during follow-up (only ICD-10 codes) among those without a diagnosis of the cancer or pre-cancer of interest at baseline. All time cancer and pre-cancer were defined as either prevalent or incident (or both) cancer and pre-cancer.

**Other covariates**

The questionnaires gave information on the covariates physical activity during leisure time (sedentary, light, or moderate/vigorous); education/vocational training (only basic education, education including students); alcohol consumption (drinks per week); smoking habits (daily smokers; or never, former and occasional smokers).
| Study population | Monica10, % (n); mean (SD) | Allergy98, % (n); mean (SD) | Inter99, % (n); mean (SD) | Health2006, % (n); mean (SD) |
|------------------|--------------------------|---------------------------|------------------------|---------------------------|
| FLG genotype     | Wild type | Mutation* | Wild type | Mutation* | Wild type | Mutation* | Wild type | Mutation* |
| Persons          | 92.5 (2,383) | 7.5 (194) | 92.9 (1,120) | 7.1 (86) | 91.2 (5,696) | 8.8 (551) | 91.1 (3,047) | 8.9 (299) |
| Age (years)      | 55.2 (10.8) | 56.1 (10.0) | 39.9 (15.1) | 42.1 (15.2) | 46.2 (7.9) | 46.5 (8.0) | 49.4 (13.0) | 49.2 (13.2) |
| P-value          | P = 0.174 | P = 0.185 | P = 0.539 | P = 0.107 |
| Male gender      | 92.8 (1,193) | 7.2 (93) | 92.9 (512) | 7.1 (39) | 91.6 (2,754) | 9.4 (285) | 91.2 (1,365) | 8.8 (132) |
| Female gender    | 92.2 (1,190) | 7.8 (101) | 92.8 (608) | 7.2 (47) | 91.7 (2,942) | 8.3 (266) | 91.0 (1,682) | 9.0 (167) |
| P-value          | P = 0.217 | P = 0.048 | P = 0.130 | P = 0.829 |
| Alcohol (drinks/week) | 10.0 (12.3) | 9.6 (12.4) | 6.8 (8.0) | 7.2 (8.9) | 10.3 (12.6) | 11.5 (19.2) | 9.6 (10.1) | 9.9 (10.9) |
| P-value          | P = 0.214 | P = 0.854 | P = 0.228 | P = 0.765 |
| Education        | 92.8 (1,778) | 7.2 (137) | 92.3 (791) | 7.7 (68) | 91.2 (4,669) | 8.8 (450) | 91.4 (2,611) | 8.6 (247) |
| No education     | 91.4 (604) | 8.6 (57) | 94.2 (335) | 5.8 (20) | 90.9 (848) | 9.1 (85) | 89.1 (392) | 10.9 (48) |
| P-value          | P = 0.217 | P = 0.247 | P = 0.782 | P = 0.121 |
| BMI (kg/m²)      | 25.9 (4.2) | 26.2 (4.1) | 25.6 (4.6) | 25.9 (4.1) | 26.3 (4.6) | 26.1 (4.5) | 25.9 (4.7) | 26.1 (5.1) |
| P-value          | P = 0.209 | P = 0.325 | P = 0.140 | P = 0.768 |
| Sedentary PA     | 93.1 (499) | 6.9 (37) | 92.3 (289) | 7.7 (24) | 91.8 (1,168) | 8.2 (105) | 89.9 (550) | 10.1 (62) |
| P-value          | P = 0.214 | P = 0.048 | P = 0.130 | P = 0.829 |
| Light PA         | 92.0 (1326) | 8.0 (115) | 92.1 (556) | 7.9 (48) | 90.8 (3479) | 9.2 (352) | 91.5 (1,838) | 8.5 (170) |
| Moderate/vigorous PA | 93.0 (518) | 7.0 (39) | 95.4 (270) | 4.6 (13) | 91.9 (949) | 8.1 (84) | 91.3 (631) | 8.7 (60) |
| P-value          | P = 0.624 | P = 0.173 | P = 0.412 | P = 0.441 |
| Never/former smokers | 91.7 (1236) | 8.3 (117) | 92.3 (741) | 7.7 (62) | 90.8 (3632) | 9.2 (369) | 91.4 (2,343) | 8.6 (220) |
| Daily smokers    | 93.3 (1,808) | 6.7 (77) | 94.0 (377) | 6.0 (24) | 91.9 (2,023) | 8.1 (179) | 89.8 (674) | 10.2 (77) |
| P-value          | P = 0.121 | P = 0.270 | P = 0.146 | P = 0.159 |
| Systolic blood pressure | 129.5 (19.3) | 127.6 (18.2) | 128.7 (17.8) | 131.8 (18.5) | 130.6 (17.6) | 130.9 (17.0) | 130.5 (17.8) | 130.7 (17.0) |
| P-value          | P = 0.215 | P = 0.058 | P = 0.541 | P = 0.685 |
| Diastolic blood pressure | 82.2 (10.5) | 81.0 (10.9) | 79.3 (11.4) | 82.1 (12.4) | 82.6 (11.4) | 82.3 (11.0) | 81.7 (10.7) | 82.9 (10.4) |
| P-value          | P = 0.095 | P = 0.070 | P = 0.954 | P = 0.059 |
| Triglycerides    | 1.5 (1.0) | 1.4 (1.0) | 1.7 (1.2) | 1.6 (0.9) | 1.3 (1.3) | 1.3 (1.3) | 1.3 (1.0) | 1.3 (0.7) |
| P-value          | P = 0.689 | P = 0.015 | P = 0.100 | P = 0.772 |
| HDL-cholesterol  | 1.4 (0.4) | 1.5 (0.5) | 1.6 (0.4) | 1.6 (0.4) | 1.4 (0.4) | 1.5 (0.4) | 1.5 (0.4) | 1.5 (0.4) |
| P-value          | P = 0.469 | P = 0.002 | P = 0.014 | P = 0.879 |

Mean (SD) and p-values from the Kruskal-Wallis test are in italic.  
* One or more FLG loss-of-function mutations  
Abbreviations: BMI, body mass index; SD, standard deviation; HDL-cholesterol, high-density lipoprotein cholesterol; FLG, filaggrin gene; PA, physical activity

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Height and weight were measured without shoes and light clothes, and body mass index (BMI) was calculated as weight divided by height squared (kg/m²). We used the average of two blood pressure measurements. In the Monica10, Inter99 and Health2006 study, serum triglycerides and HDL-cholesterol were measured from fasting blood samples using enzymatic colorimetric methods (Roche, Mannheim, Germany). In Allergy98, HDL-cholesterol and triglycerides were determined from non-fasting blood samples using the VITROS 950 automatic analyzer (Johnson & Johnson, Langhorne, Pa., USA).

Statistical analyses

The analyses were performed with SAS, version 9.2 (SAS Institute Inc. Cary, NC USA). All p-values were two-sided, and p<0.05 were considered statistically significant. Table 1 shows the baseline characteristics expressed as % (number) or mean (standard deviation, SD) according to study population and FLG genotype. Table 2 displays covariates/baseline characteristics according to all time HPV-related disease. Table 3 shows the distribution of prevalent and incident cancers and pre-cancers in the 4 studies.

Data from the four cohorts were pooled. The associations between FLG genotype and prevalent and all-time cervical cancer and pre-cancer, HPV-related cancer and all HPV-related cancer and pre-cancer were analyzed with multivariable logistic regression analyses (table 4). The estimates are presented as odds ratios (OR) and 95% confidence intervals (CI). Multivariable Cox regression analyses were used to determine the association of FLG genotype and the incidence of HPV-related cancer, cervical cancer, CIS and dysplasia, and all HPV-related cancer and pre-cancer (table 5). We used age as underlying time axis and delayed entry which means that persons enter the analysis at their age at baseline and exit the analysis at their event or censoring age. The few participants lost to follow-up (emigrated or disappeared) contributed to the risk time until the date of their last registered activity. Persons with a diagnosis of cancer and pre-cancer at baseline were excluded in the analyses of all incident HPV-related cancer and pre-cancer. Estimates are presented as hazard ratios (95% CI).

For both the logistic and Cox regression analyses, only participants with complete information on all considered variables were included. In model 1, we adjusted for gender, study population and age (not in the Cox regression analyses, since age was underlying time axis and thus accounted for). In model 2, we further adjusted for education, physical activity, smoking habits, alcohol intake, body mass index, systolic and diastolic blood pressure, serum triglycerides and HDL-cholesterol. There were no statistically significant interactions between FLG genotype and neither study population nor gender.

### Table 2. Covariates according to human papilloma virus related disease* at any time.

| Characteristics          | % (n; Mean (SD)) No HPV-related diagnosis (n = 12,790) | % (n; Mean (SD)) HPV-related diagnosis (n = 586) | P-value                          |
|--------------------------|-----------------------------------------------------|-------------------------------------------------|----------------------------------|
|                         |                                                     |                                                 | Chi-square test or Kruskal Wallis test |
| Study                   |                                                     |                                                 |                                  |
| Monica10                | 96.3 (2,481)                                        | 3.7 (96)                                        | 0.065                            |
| Inter99                 | 95.5 (5,965)                                        | 4.5 (282)                                       |                                  |
| Health2006              | 95.1 (3,181)                                        | 4.9 (165)                                       |                                  |
| Allergy98               | 96.4 (1,163)                                        | 3.6 (43)                                        |                                  |
| Gender                   |                                                     |                                                 |                                  |
| Male                    | 99.7 (6,355)                                        | 0.3 (18)                                        | <0.0001                          |
| Female                  | 91.9 (6,435)                                        | 8.1 (568)                                       |                                  |
| Age, years              | 48.3 (11.5)                                         | 47.2 (10.6)                                     | 0.026                            |
| Education               |                                                     |                                                 | 0.137                            |
| No                      | 95.0 (2,261)                                        | 5.0 (118)                                       |                                  |
| Yes                     | 95.7 (10,290)                                       | 4.3 (459)                                       |                                  |
| Body mass index, kg/m²  | 26.1 (4.5)                                          | 25.2 (5.1)                                      | <0.0001                          |
| Physical activity        |                                                     |                                                 | 0.0003                           |
| Sedentary               | 94.6 (2,585)                                        | 5.4 (149)                                       |                                  |
| Light                   | 95.6 (7,533)                                        | 4.4 (351)                                       |                                  |
| Moderate/vigorous smokers | 96.8 (2,483)                                      | 3.2 (81)                                        |                                  |
| Never/former smokers    | 96.5 (8,470)                                        | 3.5 (310)                                       |                                  |
| Daily smokers           | 93.9 (4,236)                                        | 6.1 (275)                                       | <0.0001                          |
| Alcohol, drinks/week    | 9.9 (12.1)                                          | 7.3 (8.8)                                       | <0.0001                          |
| Systolic blood pressure  | 130.4 (17.9)                                        | 125.8 (18.3)                                    | <0.0001                          |
| Diastolic blood pressure | 82.1 (11.1)                                         | 79.3 (11.2)                                     | <0.0001                          |
| Triglycerides           | 1.4 (1.2)                                           | 1.3 (0.9)                                       | 0.001                            |
| HDL-cholesterol         | 1.5 (0.4)                                           | 1.6 (0.4)                                       | <0.0001                          |

Mean (SD) and p-values from the Kruskal-Wallis test are in italic.

* Cervical cancer, carcinoma in situ and dysplasia and human papilloma virus related cancers of head and neck, vulva and vagina, penis and anus.

Mean (SD) and p-values from the Kruskal-Wallis test are in italic.

Cervical cancer, carcinoma in situ and dysplasia and human papilloma virus related cancers of head and neck, vulva and vagina, penis and anus.

Means (SD) and p-values from the Kruskal-Wallis test are in italic.

Cervical cancer, carcinoma in situ and dysplasia and human papilloma virus related cancers of head and neck, vulva and vagina, penis and anus.
Results

**FLG** mutation status was not associated with the baseline characteristics except for statistically significantly higher levels of HDL-cholesterol in the Allergy98 and the Inter99 study (table 1). Table 2 displays the covariates according to an all-time HPV-related diagnosis of cancer or pre-cancer. As expected due to the contribution of cervical cancers and pre-cancers, a HPV-related diagnosis is significantly associated with female gender in crude analyses. In addition and also in crude analyses, an all-time HPV-related diagnosis is significantly associated with younger age, lower BMI, daily smoking, lower alcohol consumption, lower systolic and diastolic blood pressure, lower triglycerides and higher HDL-cholesterol (table 2).

Table 3 shows the distribution of prevalent and incident cancers and pre-cancers in the 4 studies. For HPV-related cancer in particular, the number of events in some of the categories is very low (table 3). In table 4, the associations between **FLG** genotype and prevalent and all-time HPV-related cancers and pre-cancer are shown. With a total of 534 events, the association between **FLG** genotype and all HPV-related disease (all-time) was statistically significant with hazard ratio, HR = 2.0 (95% CI: 1.0, 1.8) for **FLG** wild type in the fully adjusted model (table 3, model 2). In general, the associations remained essentially unchanged by multiple adjustments.

Discussion

**FLG** mutation carrier status was significantly associated with a higher risk of incident cervical cancer and pre-cancer and all HPV-related cancers and pre-cancers in Danish adults. Furthermore, a significantly higher risk was found among **FLG** mutation carriers of prevalent or incident HPV-related cancer or pre-cancer. To our knowledge, no study has investigated this before. Although previous histological studies on filaggrin and the cervix aimed to evaluate the use of filaggrin expression as a diagnostic criteria in cervical lesion, the conclusions are somewhat in line with our results: Cintorino et al found that filaggrin expression was more irregular in the high risk HPV type cervical lesions (HPV 16 and 18) as compared to the low risk (HPV 6, 11 and 31) [25]. Lara et al found that filaggrin expression could serve as a marker of differentiation in both normal and pathological cervical tissue and that even neoplastic lesion may have regular filaggrin expression if well differentiated [26]. Thus, the above-mentioned studies suggest that disturbed filaggrin expression is more often seen in less differentiated pre-cancers and cancers.

**FLG** mutation carrier status could affect the risk of HPV-related cancer and pre-cancer in several ways. First, since the skin is an important barrier against microorganisms, impaired skin barrier function caused by filaggrin deficiency may lead to a greater susceptibility to microorganisms [7]. Hence, Milden et al found that mutation carriers had impaired filament aggregation and a reduced number of tight junctions in a skin model [11]. The absence of filaggrin resulted in a higher UV sensitivity, likely due to a smaller amount of epidermal urocanic acid, a breakdown product of the filaggrin molecule. As a possible consequence, **FLG** mutations are associated with up to 10% higher levels of vitamin D, possibly due to higher UV sensitivity [27,28]. Also, a

### Table 3. Distribution of prevalent and incident human papilloma virus related disease according to study population and **FLG** genotype.

| **FLG** genotype | HPV-related cancer*, % (n/n_{total})** | Cervical cancer, CIS and dysplasia***, % (n/n_{total})**** |
|------------------|----------------------------------------|----------------------------------------------------------|
|                  | Wild type | Loss-of-function mutation | Wild type | Loss-of-function mutation |
| Total            | Prevalent | 0.3 (33/12246) | 0.1 (1/1130) | 6.8 (438/6422) | 7.9 (46/581) |
|                  | Incident | 0.2 (21/12246) | 0.4 (4/1130) | 1.0 (62/6422) | 1.7 (10/581) |
| Monica10         | Prevalent | 0.3 (7/2383) | 0.5 (1/194) | 6.1 (72/1190) | 7.9 (8/101) |
|                  | Incident | 0.3 (7/2383) | 1.0 (2/194) | 0.6 (7/1190) | 1.0 (1/101) |
| Allergy98        | Prevalent | 0.4 (4/1120) | 0 (0/86) | 3.8 (23/608) | 4.3 (2/47) |
|                  | Incident | 0.1 (1/1120) | 0 (0/86) | 2.0 (12/608) | 4.3 (2/47) |
| Inter99          | Prevalent | 0.2 (13/5696) | 0 (0/551) | 7.4 (219/2942) | 8.6 (23/266) |
|                  | Incident | 0.1 (8/5696) | 0.2 (1/551) | 0.9 (26/2942) | 1.5 (4/266) |
| Health2006       | Prevalent | 0.3 (9/3047) | 0 (0/299) | 7.4 (124/1682) | 7.8 (13/167) |
|                  | Incident | 0.2 (5/3047) | 0.3 (1/299) | 1.0 (17/1682) | 1.8 (3/167) |

* HPV-related cancers include cancer, cervix (ICD-7: 1711, 1712, ICD-10: C53), cancer, head and neck, subset (ICD-7: 1410, 1411, 1418, 1450, 1442, 1458, 1480, ICD-10: C01.9, C02.4, C02.8, C09.0, C09.1, C09.8, C09.9, C10.2, C10.8, C10.9, C14.0, C14.2, C14.8), cancer, vulva and vagina (ICD-7: 1760, ICD-10: C51, C52), cancer, penis (ICD-7: 1790, ICD-10: C60), and cancer, anus (ICD-7: No sufficiently specific ICD-7 code, ICD-10: C21.0-C21.8).

** Of the whole population, i.e. both men and women

*** Cervical cancer, CIS and dysplasia include cancer, cervix (ICD-7: 1711, 1712, ICD-10: C53), CIS, cervix (ICD-7: 5710, 5711, ICD-10: D06), and dysplasia, cervix (ICD-7: 5712, ICD-10: N87).

**** Of women

Abbreviations: CIS, carcinoma in situ; HPV, human papillomavirus; **FLG**, filaggrin gene

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breakdown products of filaggrin to slow the growth of microorganisms [7]. Thus, Miajlovic et al found the principal degradation products of filaggrin may lead to increased adhesion of Staphylococcus aureus. **Molluscum virus** [30].

Cutaneous viral infections such as herpes simplex virus, HPV, or **frequently complicated by both localized and disseminated infection** [7]: the skin of persons with atopic dermatitis are susceptible to cutaneous microorganism colonization and is characterized by a skin barrier defect and an increased susceptibility to cutaneous microorganism colonization and infection [7]: the skin of persons with atopic dermatitis are frequently colonized by S. aureus, and atopic dermatitis is rather frequently complicated by both localized and disseminated cutaneous viral infections such as herpes simplex virus, HPV, or molluscum virus [30].

Second, elevated pH of the stratum corneum due to less acidic lipophilic permeability of stratum corneum in a skin equivalent in a skin equivalent [29]. They concluded that FLG knockdown alone may not necessarily affect the skin barrier function [29]. However, atopic dermatitis affects approximately 40% of FLG mutation carriers who have less FLG loss-of-function mutation carriers with atopic dermatitis [31]. In addition, Gao et al found a higher risk of eczema herpeticum in FLG mutation carriers with atopic dermatitis compared with FLG wild types with atopic dermatitis [32].

Third, a low grade skin inflammation can promote the conditions for neoplastic cells to proliferate thus increasing the risk of local cancer [12-14]: inflammation contributes to proliferation and survival of malignant cells, angiogenesis and metastasis, and induction of genetic instability with accumulated random genetic changes in cancer cells [13].

Further research into a potential effect of FLG loss-of-function mutations on other microorganism related cancers is important. Other bacteria and viruses known to be carcinogenic are hepatitis B and C virus (hepatocarcinoma); Epstein-Barr virus (lymphoma and nasopharyngeal carcinoma); and helicobacter pylori (gastric cancer) [12,33]. Also, suspected to be carcinogenic are hepatitis C virus (hepatocarcinoma), streptococcus bovis (cancer of the gallbladder), and chlamydia pneumonia (lung cancer) [12,33].

The strengths of our study include the prospective design and the large population-based samples; a long-term follow-up and the use of standardised registry-based diagnoses with a high degree of

### Table 4. Odds ratios and 95% confidence intervals for the associations between FLG genotype and prevalent and all time HPV-related disease.

| Prevalent disease | Events (persons included) | Model 1 OR (95% CI) | Model 2 OR (95% CI) |
|-------------------|---------------------------|---------------------|---------------------|
| HPV-related cancer* | 33 (12,442) | 0.3 (0.05, 2.4) | 0.3 (0.04, 2.4) |
| FLG loss-of-function mutation vs. wild type | | P = 0.274 | P = 0.269 |
| Cervical cancer, CIS and dysplasia*** | 437 (6,417) | 1.2 (0.9, 1.7) | 1.2 (0.9, 1.7) |
| FLG loss-of-function mutation vs. wild type | | P = 0.285 | P = 0.256 |
| All HPV-related disease*** | 442 (12,442) | 1.2 (0.9, 1.6) | 1.2 (0.9, 1.7) |
| FLG loss-of-function mutation vs. wild type | | P = 0.324 | P = 0.294 |

| All time disease (prevalent and incident) | | | |
|-------------------|---------------------------|---------------------|---------------------|
| HPV-related cancer* | 55 (12,442) | 1.1 (0.4, 2.7) | 1.1 (0.4, 2.7) |
| FLG loss-of-function mutation vs. wild type | | P = 0.875 | P = 0.885 |
| Cervical cancer, CIS and dysplasia*** | 506 (6,417) | 1.3 (1.0, 1.8) | 1.3 (0.98, 1.8) |
| FLG loss-of-function mutation vs. wild type | | P = 0.081 | P = 0.068 |
| All HPV-related disease*** | 534 (12,442) | 1.3 (1.0, 1.8) | 1.4 (1.0, 1.8) |
| FLG loss-of-function mutation vs. wild type | | P = 0.050 | P = 0.040 |

Abbreviations: CI, confidence interval; OR, odds ratio, CIS, carcinoma in situ; HPV, human papillomavirus.

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hydrophilic fluorescent dye easily penetrated through the stratum corneum down to the basal layer of the filaggrin-deficient skin cultures [11]. Of note, a recent study found that filaggrin knockdown did not affect either epidermal morphogenesis, lipid organization in stratum corneum, lipid composition, or the lipophilic permeability of stratum corneum in a skin equivalent [29]. They concluded that FLG knockdown alone may not necessarily affect the skin barrier function [29]. However, atopic dermatitis affects approximately 40% of FLG mutation carriers who have less FLG loss-of-function mutation carriers with atopic dermatitis [31]. In addition, Gao et al found a higher risk of eczema herpeticum in FLG mutation carriers with atopic dermatitis compared with FLG wild types with atopic dermatitis [32].

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Further research into a potential effect of FLG loss-of-function mutations on other microorganism related cancers is important. Other bacteria and viruses known to be carcinogenic are hepatitis B and C virus (hepatocarcinoma); Epstein-Barr virus (lymphoma and nasopharyngeal carcinoma); and helicobacter pylori (gastric cancer) [12,33]. Also, suspected to be carcinogenic are salmonella typhi (carcinoma of the gallbladder), streptococcus bovis (colorectal cancer), and chlamydia pneumonia (lung cancer) [12,33].

The strengths of our study include the prospective design and the large population-based samples; a long-term follow-up and the use of standardised registry-based diagnoses with a high degree of
suggesting that the observed association between related to several factors that may be related to cancer risk exposure happens before the outcome, eliminating the risk of pleiotropic effects of HPV caused by HPV differs between the included cancer status was not associated with mortality [20]. Also, the proportion completeness and a minimal loss to follow-up. Using a genetic marker such as also establishes the time sequence i.e. that the exposure happens before the outcome, eliminating the risk of reverse causation. As shown in table 1, FLG mutations were not related to several factors that may be related to cancer risk suggesting that the observed association between FLG mutations and HPV-related cancer and pre-cancer risk is not mediated by pleiotropic effects of FLG mutations on these factors. The validity of the cancer diagnoses in the Danish Cancer Registry is high with the proportion of morphologically verified tumors of 89% [21], a validity secured through daily quality control routines and in completing the yearly publication of the Cancer Registry and the quality achieved by e.g. manual coding of complex cases [21]. HPV is considered mandatory for cervical cancer development whereas the fraction of cancers caused by HPV differ among other HPV-related cancers; possible delay from onset of disease until inclusion in the register; and the risk of selection bias/survivor bias due to FLG deficiency that leads to increased retention and penetration of HPV in tissue with eventual cancer development.

## Author Contributions

Conceived and designed the experiments: TS LLNH AL TJ JDJ JPT SS PBS TM PB. Performed the experiments: AL TJ JDJ JPT SS PBS TM. Analyzed the data: TS LLNH AL. Contributed reagents/materials/analysis tools: AL TJ. Wrote the paper: TS AL LLNH.

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## Table 5. Hazard ratios and 95% confidence intervals for the associations between FLG genotype and incident HPV-related disease.

| FLG genotype                  | Events (persons included) | Model 1 HR (95% CI) | Model 2 HR (95% CI) |
|-------------------------------|---------------------------|---------------------|---------------------|
| FLG wild type                 | 22 (12,409)               | 1 (reference)       | 1 (reference)       |
| FLG loss-of-function mutation | 2.4 (0.8, 7.0)            | P = 0.114           | P = 0.122           |
| Cervical cancer, CIS and dysplasia | 69 (5,980) | 2.0 (1.0, 3.8)      | P = 0.050           | 2.0 (1.0, 4.0)      | P = 0.038           |
| FLG wild type                 | 1 (reference)             |                      | 1 (reference)       |
| FLG loss-of-function mutation | 2.1 (1.2, 3.7)            | P = 0.014           |                    |
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