Incidence of IP and risk of malignant transformation in the Swedish population 1960–2010

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Abstract The true incidence of inverted papilloma (IP) is not yet known. From hospital-based studies, its incidence has been estimated to approximately 0.5/100,000 person years. Earlier hospital case studies have shown that IP can undergo a malignant transformation in 1–53%. The frequency of its malignant transformation on a population basis is unknown. To our knowledge, no standardised incidence ratio (SIR) has been reported for malignancies among IPs. This study aims to investigate these incidences on a population basis. Using the data from the Swedish Cancer Registry (SCR), we have identified patients with IP and patients with Squamous Cell Carcinoma (SCC) diagnosed between 1960 and 2010 in Sweden. Incidence of IP and incidence of SCC among patients with IP and SIR were analyzed. Eight hundred and fourteen patients with IP were identified. The incidence of IPs reported to the SCR increased from 1960 to 2010. In this cohort, SCC was overrepresented, as compared with the general population. The incidence of IP in the Swedish population seems to have increased.

Keywords Inverted papilloma · Squamous cell carcinoma · Neoplastic cell transformation · Incidence · Registries

Introduction

Inverted papilloma (IP), often referred to as schneiderian papilloma, is a locally destructive benign tumour of the sino-nasal mucosa with a tendency for malignant transformation and a high propensity for recurrence [1–3]. It arises from the transitional epithelium, the Schneiderian membrane. Histologically, IP typically comprises both exophytic and endophytic components. The tumour invaginates into the underlying tissue, hence its name [4].

The true incidence of IP is not yet known, nor is its incidence pattern over time. From hospital-based studies, the incidence has been estimated to approximately 0.5/100,000 person years [1, 5, 6]. Most patients are diagnosed in their 60s with a male predominance of 2–10:1 [7, 8]. Earlier hospital case studies have shown that IP undergoes a malignant transformation in 1–53% [9, 10]. The frequency of its malignant transformation on a population basis is unknown. To our knowledge, no standardised incidence ratio (SIR) has been reported for malignancies among IPs. The malignancies can occur synchronously, diagnosed in the same initial lesion, or metachronously, arising in the same area where the IP had previously been removed. In the literature, synchronous cancers are more frequent [11].

The aim of the present population-based cohort study was to establish the incidence of IP over time in the Swedish population as well as the risk of concomitant and later malignancies in the sino-nasal mucosa in patients with IP.

Materials and methods

Subjects

The Swedish Cancer Registry (SCR) covers the whole Swedish population, today approximately nine million
people. All malignant tumours as well as some precan-
cerous tumours, comprising IP, are reported to the SCR
since 1958. The reliability of the SCR is high and the
reporting to SCR is estimated to over 96 %, at least
regarding true malignant neoplasms [12, 13]. Approxi-
mately, 99 % of the cases are morphologically verified
[12]. The SCR reports cases using the ICD-7. Clinical
diagnoses according to ICD-8 to ICD-10 have been trans-
fomed to ICD-7 in the register.

From the SCR, we included all tumours in the sino-nasal
area, numbered 160.0, 160.2, and 160.7–160.9, according
to the ICD-7, with a histological code for true papillomas.
We chose to include patients reported to the SCR with IP
since 1960, since the reporting to the register is considered
reliable since 1960 [13]. Through the data from SCR, we
were able to identify other information, such as gender
distribution and age, at diagnosis (Table 1). In a similar
way, we also retrieved data about those patients who were
diagnosed with an SCC in the sino-nasal mucosa.

Statistics

Using the emigration and immigration registers from
Statistics Sweden, we were able to identify patients lost to
follow-up. Data concerning the size of the population and
its age and sex distributions from year to year were also
given by Statistics Sweden.

Because of the rarity of IP, we chose to report its inci-
dence per decade. The calendar period was divided into
decades beginning in 1960. Incidence rates were calculated
by dividing the number of cases in each calendar period by
the total average population in each age group in respective
calendar period. The incidence rates were calculated for the
whole population as well as sex specific. For comparison,
the rates are also age-adjusted to the Swedish Standard
Population of year 2000.

| Table 1 | Characteristics of patients with IP and SCC in Sweden 1960–2010 |
|---------|---------------------------------------------------------------|
|         | 1960’s | 1970’s | 1980’s | 1990’s | 2000’s | Total |
| IP      | N men  | 3      | 36     | 165    | 148    | 232    | 584    |
|         | Mean age | 40.7   | 57.6   | 52     | 54.9   | 56.3   | 54.7   |
|         | N women | 3      | 15     | 69     | 48     | 95     | 230    |
|         | Mean age | 47.7   | 54     | 55.9   | 52     | 59     | 56.1   |
|         | N total | 6      | 51     | 234    | 196    | 327    | 814    |
|         | Mean age | 44.2   | 56.5   | 53.1   | 54.2   | 57.1   | 55.1   |
|         | Men/women | 1      | 2.4    | 2.4    | 3.1    | 2.5    | 2.5    |
| SCC     | Total   | 1      | 4      | 6      | 11     |        |
|         | Mean age | 66     | 69.5   | 61.2   | 64.6   |

The risk of developing cancer among patients with IP
was calculated as a proportion and a standardised incidence
ratio (SIR). SIRs were calculated for SCC in patients
diagnosed with IP by dividing the observed numbers of
SCC in this group by the expected numbers of SCC based
on person years at risk and population incidence.

Statistical analysis was done in SAS 9.2, STATA, and
Excel.

The study was approved by the Ethical Committee at the
Karolinska Institute, Stockholm, Sweden, according to the
ethical permission 2012/49-31/2.

Results

The mean age of diagnosis for IP was 55.1 (54.7 in male
patients and 56.1 for women). No trend was seen over time
(Table 1).

The male-to-female ratio in the 60s was 1:1, but only six
cases were reported. The male-to-female ratio was rather
constant over time from the 70s, 2.4–3.3:1 with no time-
dependent trend. The incidence of IP increased over time
with a substantial leap from the 60s when the incidence
was 0.01/100,000 py (person years), to the 70s when the
incidence was 0.07/100,000 py, and from the 70s to the 80s
when the incidence was 0.29/100,000 py. After a smaller
decrease during the 90s, \( i = 0.23/100,000 \) py, the inci-
dence has again increased during the first decade of the
twenty-first century to 0.33/100,000 py (Fig. 1). The same
trend was seen in male and females.

Of all the persons who were diagnosed with IP, \( n = 814, 11 \) were diagnosed with SCC or SCC in situ
which corresponds to a proportion of 1.35 % (Table 2).
This represents an incidence of SCC over the whole time
period of 111/100,000 py among patients with IP as com-
pared with 0.37/100,000 py in the general population. Only
in the male population did the IPs undergo a malignant
transformation, which is a statistically significant
(p < 0.05) difference compared with women. Among men with IP, the proportion developing SCC was 1.88 % with an incidence of 157.8/100,000 py as compared with 0.46/100,000 py in the general male population.

For the whole IP cohort, SIR was 142.76 with a p value <0.05, thereby also showing an over-representation of SCC among patients with IP. When analyzing the SIR stratified by duration, we found no evidence of reverse causality (Table 3).

We did not find that patients with IP had a higher risk of developing sino-nasal malignancies of other histological types than SCC compared with the general population (data not shown).

### Discussion

In this population-based study, we found that during the study period, the age standardised incidence of IP increased by approximately 400 %, from 0.01 to 0.33/100,000 py.

The SIR of 142.76 showed a large over-representation of SCC among patients with IP (p < 0.005). Interestingly, the malignant transformation only occurred in men, representing a proportion of 1.88 % among men with IP. This represents a 300-fold increased risk as compared with the general male population.

In our study, two persons (18 %) with IP had synchronous SCC and nine (82 %) had metachronous SCC. Although our study was population based, our results regarding the incidence of IP are well in line with hospital data presented in Denmark [1, 14] and in the European position paper on the endoscopic management of tumours of the nose, paranasal sinuses, and skull base conducted by Lund et al. [15].

There are, however, some differences, where we found an age standardised incidence of 0.29 in the 80’s, whereas in Denmark, the incidence was 0.52 during approximately the same time period. This could be due to hazard but may also show the advantage of a higher coverage in a population-based study, or less accurate histopathological diagnosis.

In contrast to malignant sino-nasal tumours (malignant melanoma excluded) which have decreased over the decades [16], we found an increase of IP over time: In the 60’s, the incidence was very low (0.008), but from the 70’s, there has been a fivefold increase (0.006–0.033), an increase which has not been shown earlier. This may be a true increase. However, this could very possibly be due to underreporting during the earlier part of the study period and the possibility that over time more sino-nasal biopsies were sent for pathological assessment and diagnosis. The accuracy in the Swedish Cancer registry has been investigated for common, malignant tumours. IP is considered a precancerous lesion and the reporting rate may, therefore, be lower than for true malignant lesions, especially in the beginning of the study period. The increase is possibly also related to the detecting of IP due to a higher awareness of IP among ENT physicians and in their use of endoscopic sinus surgery. The preciseness of the histopathological evaluation has improved and may have lead to an increased reporting to the cancer register. The increase may also be correlated to a, until now, not fully identified environmental factor.

Earlier studies have reported diverging results concerning the risk of malignant transformation of 1–53 %, most of which are considerably higher than the 1.35 % that we found in our population-based study [8–10, 17, 18]. This discrepancy may be due to that earlier hospital-based studies from referral centres have dealt with patients with larger or recurrent IP tumours and that earlier studies, not based on a register with complete coverage of the population, (such as the SCR), have not included all IPs. It may also be due to differences in studied populations. Another plausible cause for the higher risk of malignant transformation in the previous studies is the likely underreporting of well-differentiated SCC “ab initio”. However, percentage does not allow comparison between studies on different populations, since difference in age or gender distribution is not known. Hopefully, future studies will be conducted presenting SIRs in other regions allowing a better comparison.

### Table 2 Sex distribution of patients with IP and, among them, with SCC in Sweden 1960–2010

| Sex     | N women | N men |
|---------|---------|-------|
| IP      | 230     | 589   |
| SCC     | 0       | 11    |

| N women number of women diagnosed with IP and among them SCC, N men number of men diagnosed with IP and among them SCC |

### Table 3 SIR of SCC in patients with IP stratified by duration of IP

| N | Duration | Expected | Outcome | SIR | SIRL | SIRU | p    |
|---|----------|----------|---------|-----|------|------|------|
| 1 | 3–4 years| 0.013621 | 1       | 73.42| 1.86 | 409.06| 0.0773|
| 2 | 5–9 years| 0.019131 | 3       | 156.81| 32.34 | 458.28| 0.0014|
| 3 | More than 10 years | 0.037111 | 4       | 107.78| 29.37 | 275.97| 0.0003|

N number of cases with IP who develop SCC, duration time between diagnosis of IP and of SCC, expected expected cases of SCC, outcome accumulated number of cases, SIR standardised incidence ratio, SIRL SIR lower, SIRU SIR upper, p p value
Our finding that only men with IP developed SCC has not been described earlier. To verify if this discrepancy between genders is true, larger cohort studies are needed. When considering the risk of SCC in the whole IP cohort, we should have found approximately five cases of SCC in the female population. In the review of the English literature conducted by Nudell et al. [10], 11 out of 59 cases of SCC ex IP occurred in women.

In the literature, synchronous SCC has previously been reported to be more common than metachronous malignant transformation [19]. In a meta-analysis including more than 2000 cases from 2007, Mirza and co-workers found a risk of 7.1% of synchronous carcinomas compared with a risk of 3.6% of metachronous malignancies [11]. Interestingly in our study, metachronous cancers were more than four times as common as the synchronous ones. A similar proportion was seen in only 4 of the 64 studies in the meta-analysis. Differing rates found in these earlier studies may be exaggerated due to referral bias to tertiary centres. Again, the difference in findings could be due to underreporting by the pathologist of IP when SCC is found in the same specimen. This needs to be investigated further.

Conclusion

In this first population-based study, the incidence of IP has increased from 0.01 to 0.33/100,000 py in Sweden from 1960 to 2010. The risk of SCC malignancies in the IP cohort is lower than in the previous studies although significantly higher than in the general population. Interestingly, we only found SCC among male patients with IP where the risk of malignant transformation represented a 300-fold increased risk compared with the general male population.

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Compliance with ethical standards

Conflict of interest No conflicts of interest for any of the authors.

Ethical approval The study has been approved by the Regional Ethics Committee at the Karolinska Institute. This article does not contain any studies with human participants or animals performed by any of the authors.

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