The health effects derived from UV radiation and sunbed use

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Outline of this presentation

• UV and skin cancer
• Sunbed use – who and why?
• Tanning – is it worth it?
• Sunbed addiction
UVB: 280-320 nm

- Skin tanning, erythema formation (snow blindness in eye)
- A Full Carcinogen (both promoter and initiator)
- Causes DNA damage directly -> leads to UV-signature mutations (CC to TT transitions)
- Immunosuppressive
- Vitamin D synthesis

UVA: 320- 400 nm

- Less carcinogenic than UVB; DNA damage mainly via reactive oxygen species (ROS)
- Photoaging
- Immunosuppressive
UV radiation as a risk factor for skin cancer formation

- Skin cancer incidence has increased substantially over the past decades and the role of UV radiation in the etiology of skin cancer is well established.

- Basal cell carcinoma (BCC), and especially Squamous cell carcinoma (SCC), are related to the cumulative, lifetime UV exposure.

- Both types are prevalent at the elderly age groups on the sun exposed areas.

 Basal cell carcinoma (8528 cases in 2016)  
 Squamous cell carcinoma (1719 cases in 2016)
Malignant melanoma is one of the fastest increasing cancer in incidence in the western world

- Associated strongly with intermittent UV exposure (history of sunburns), rather than chronic exposure, with genetically susceptible individuals
- Intermittent exposure in childhood and adolescence has an important role, although the adult exposure contributes as well
- Prevalent among younger adults as compared to BCC and SCC

- There were 1947 melanomas in Finland in 2016
- There were 212 melanoma deaths in Finland in 2016
Sunbeds now and then

• Sunbeds were first introduced to the market in the late 1970’s\(^1\)
  – Mercury lamps emitted mainly UVB wavelengths and even some UVC
  – Mainly used at homes

• In 1980’s indoor tanning became more popular in beauty salons\(^1\)
  – Lamps emitted 99% of UVA and 1% of UVB

• Nowadays sunbeds contain as much UVB as in the sunlight, but **considerably more UVA as compared to the sun**\(^2,3\)

• **Repeated exposure to large amounts of UVA** in relatively short periods (10-20 mins per tanning session) **constitutes a new experience for skin**

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3 Nilsen et al. BJD. 2016; 2016; 174: 730-740
Sunbed power indicated by UV index*

- Sun, Helsinki Finland, December 0.1
- Sun, Helsinki Finland, Mid-Summer 6-7
- Sun, Mediterranean, Mid-Summer 11
- Sunbeds in EU 12
- Sunbeds out of EU even up to 24

- Phototherapy devices in hospitals
  - PUVA (Psoralen+UVA) 4 - 7
  - SUP (315-360 nm) 20 - 32
  - Narrow-band UVB (309-313 nm) 210 - 240

*UV index is calculated multiplying erythema-weighted irradiance by 40
Sunbed use – who and why

- Indoor tanning is a widespread practice in most western countries, particularly in Northern Europe and the USA
- The typical sunbed user is a female, between 17-35 y old (Schneider and Krämer, 2010)

<table>
<thead>
<tr>
<th>Reason</th>
<th>1997*</th>
<th>2011*</th>
<th>2016*</th>
<th>2018*</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cosmetic Tanning</td>
<td>30%</td>
<td>44%</td>
<td>76 (-)%</td>
<td>23 (25)%</td>
</tr>
<tr>
<td>Tanning before a trip</td>
<td>36%</td>
<td>28%</td>
<td>6 (18)%</td>
<td>58 (10)%</td>
</tr>
<tr>
<td>Doctor’s referral</td>
<td>4%</td>
<td>5%</td>
<td>2 (2)%</td>
<td>- (-)</td>
</tr>
<tr>
<td>‘Self-medication’</td>
<td>8%</td>
<td>4%</td>
<td>1 (28)%</td>
<td>3 (37)%</td>
</tr>
<tr>
<td>Mood uplift</td>
<td>11%</td>
<td>3%</td>
<td>- (30)%</td>
<td>5 (18)%</td>
</tr>
<tr>
<td>General health uplift</td>
<td>-</td>
<td>2%</td>
<td>14 (7)%</td>
<td>10 (-)%</td>
</tr>
</tbody>
</table>

*Survey performed by 1997 by V. Jalarvo, GraduThesis
*Survey performed in 2011 by Taloustutkimus Oy
*Survey performed in 2016 by Taloustutkimus Oy
*Survey performed in 2018 by Taloustutkimus Oy
Tan or pigmentation on skin – is it protective?
Yes and no….

• Constitutive epidermal melanin seems to be rather effective in darker skin types to protect the underlying cells from UV radiation\textsuperscript{1,2}
  
  – Provides sun protection factor (SPF) \textasciitilde 13
  – Pretty good ‘real’ protection for skin cells in UV-independent process

• Fair skin types react to UV radiation by facultative pigmentation, i.e tanning
  – Provides sun protection factor (SPF) \textasciitilde 2-3

\textsuperscript{1} Tadokoro et al. J Invest Dermatol. 2005; 124: 1326-1332
\textsuperscript{2} Coelho and Hearing. Pigment Cell Melanoma Res. 2010; 23(1): 57-63
Sunbed-derived tanning – is it worth it?

• UVA-rich tanning lamps have previously been promoted as ‘damage-free’, however this message could be a deceptive one
  – UVA does not increase melanin production or redistribution in epidermis, mainly photo-oxidation (darkening) of the existing tan
  – UVA-induced tan offers no photo-protection against UV exposure

• UVA is a possible carcinogen, especially for melanoma formation
  – Several animal models (fish, opossum, mice) suggest that UVA might have a role in melanomagenesis

• Sunbed is not an optimal way to ensure vitamin D synthesis due to the unnaturally large amount of UVA radiation that comes along
Sunbed use and skin cancer

- International Agency for Research on Cancer (IARC) classified artificial UV tanning devices as carcinogenic to humans

  - A ‘hall mark’ meta-analysis found a significant increase in risk of malignant melanoma, especially if sunbed use started before age of 35 (IARC 2006)
  - Epidemiological data published after the original IARC report strengthens the link between MM and artificial tanning 1-3
  - 7% of malignant melanomas (MM) in women, and 4% of MM in men, has been estimated to be related to the sunbed use4
  - Sunbed use has been also linked to SCC and BCC formation, especially if sunbed use started before age of 25 years5

- Many countries have passed laws, that restrict the sunbed use for minors (18 y) to protect young people from the UV-derived health hazards

1 Lazovich et al. Cancer Epidemiol Biomarkers Prev. 2010; 19(6)
2 Veierod et al. Cancer Epidemiol Biomarkers Prev. 2010; 19(1)
4 Boniol et al. BMJ. 2012; 345
5 Wehner et al. BMJ. 2012; 345
Sunbed addiction

Several studies have suggested that tanning behavior exhibits signs of psychologic and physiologic dependence

− First postulated in 1983, when UVA (but not visible light) increased a elevation of plasma endogenous opioid levels\(^1\)
− Surveys studies have reported of relaxation, pain relief and positive mood effects\(^2,3\)

− In an experimental sham study (blinded trial), frequent solarium users seemed to sense with their skin the true solarium from a sham device\(^4\)

− A subsequent study in 2006 showed that an opioid blockade (by naltrexone) induced withdrawal symptoms for frequent tanners\(^5\)

\(^1\) Levins et al. Lancet 1983; 2:166
\(^3\) Kaur et al. Photochem Photobiol Photodermatol 2005
Mechanism for action – a hypothesis

- UV-derived DNA damage activates the tumor suppressor protein p53 (guardian of genome) in keratinocytes

- p53 stimulates the transcription of pro-opiomelanocortin (POMC)

- Cleavage products of POMC are Melanocyte-Stimulating Hormone (MSH-α) and β-endorphin

- MSH activates the melanin synthesis in melanocytes (tanning)

- β-endorphin expression may be in response in reinforcing UV-seeking behavior of heavy tanners

Fisher and James, NEJM, 2010; 363:903-903
Working model of endogenous opioid dependence caused by chronic UV exposure

UV induces p53 signalling in keratinocytes, increasing the synthesis of POMC peptide and the concomitant β-Endorphin in the skin, leading eventually to the elevated plasma levels of β-Endorphin.

Sustained levels of plasma β-Endorphin, increase signaling at opioid receptors in the central nervous system, producing the endogenous opioid-dependent state.

Tejeda and Bonci, Cell, 2014; 157: 1500-1501
“Take home messages”

• There is no safe tan from natural or artificial UV-sources

• Sunbed use has been linked with skin cancer formation, especially with melanoma

• Solarium use restrictions aim to protect adolescents from UV derived health problems and also to minimize health care expenses for the long run

• Ultraviolet tanning can be addictive through β-Endorphin secretion from skin cells after UV exposure