Expression of individual lamins in basal cell carcinomas of the skin

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Summary In this study we used a unique collection of type specific anti-lamin antibodies to study lamin expression patterns in normal human skin and in skin derived from patients with basal cell carcinomas (BCCs). Lamin expression in serial sections from frozen tissue samples was investigated by single and double indirect immunofluorescence. In normal skin, lamin A was expressed in dermal fibroblasts and in suprabasal epithelial cells but was absent from all basal epithelial cells. Lamin C was expressed in dermal fibroblasts, suprabasal epithelial cells and a majority of basal epithelial cells. However, lamin C was not expressed in quiescent basal epithelial cells. Lamin B1 was expressed in all epithelial cells but was not expressed in dermal fibroblasts. Finally, lamin B2 was expressed in all cell types in normal skin. Lamin expression was also investigated in a collection of 16 BCCs taken from a variety of body sites. Based upon patterns of lamin expression the BCCs were classified into four groups: A-negative (10/16 tumours), C-negative (5/16 tumours), A/C-negative (1/16 tumours) and A/B2-negative (1/16 tumours). Lamin expression was also compared to cell proliferation index by staining serial sections with the proliferation marker Ki67. 9/10 of the lamin A negative tumours were highly proliferative, whereas 4/5 of the lamin C negative tumours were slow growing. Thus as a general rule absence of lamin A was correlated with rapid growth within the tumour, while absence of lamin C was correlated with slow growth within the tumour. Our data supports the hypothesis that lamin A has a negative influence on cell proliferation and its down regulation may be a requisite of tumour progression.

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The nuclear lamina is a meshwork of intermediate filaments that lines the nucloplasmic face of the nucleus and interacts with nuclear pores (reviewed by Hozak et al, 1995; Vaughan et al, 2000). Lamins, or type V intermediate filaments, consist of two subfamilies termed A-type and B-type. Whilst A and C lamins (the A-type subfamily) are alternative splice products of a single gene, there are at least four B-type genes in vertebrates (lamins B1, B2, B3 and IV (Benavente et al, 1985a; Quinlan et al, 1995). Lamins have been described as a number of diverse functions including roles in the targeting of nuclear membrane vesicles to chromatin after mitosis (Burke and Gerace, 1986), correct assembly of nuclear pores (Goldberg et al, 1995) and indirectly in the organization of replication domains (Jenkins et al, 1993; Zhang et al, 1996).

A-type and B-type lamins differ in several respects including their behaviour at mitosis (Gerace and Blobel, 1980), their post-translational processing (Farnsworth et al, 1989) and their expression during differentiation. B-type lamins are ubiquitous components of all cells, although different cells do express different B-type lamins (Stick and Hausen, 1985; Furukawa and Hotta, 1993). The two most prominent B-type lamins in mammalian somatic cells are lamins B1 and B2, although not all somatic tissues express both of these B-type lamins (Benavente and Krohne, 1985b; Stick and Hausen, 1985; Lehner et al, 1987; Broers et al, 1997).

The expression of A-type lamins appears to be linked to differentiation and is first observed in embryos at the time of organogenesis (Schatten et al, 1985; Stewart and Burke, 1987; Röber et al, 1989). Pluripotent cells can be induced to express A-type lamins in culture by treatment that induces differentiation (Lebel et al, 1987) and in some instances the ectopic expression of lamin A can promote differentiation (Lourim and Lin, 1992). The correlation between A-type lamin expression and cell differentiation has led some workers to speculate that A-type lamins may facilitate differential gene expression, by anchoring chromatin at the nuclear envelope or by sequestering inhibitors (Nigg, 1989). Consistent with this hypothesis, lamin A displays a higher affinity for chromatin binding than B-type lamins (Höger et al, 1991). A-type lamins also bind the negative growth regulator p110(R) (Ozaki et al, 1994).

The expression of A-type lamins is also altered in many tumours. In lung tumours there is an inverse correlation between the level of the proliferation marker Ki67 staining with that of the anti-lamin A/C monoclonal BU31 (Rowlands et al, 1994). Changes in the expression of lamin A has also been reported in testicular germ cell tumours (Machiels et al, 1997) and Hodgkin's disease (Jansen et al, 1997). In addition, adrenal cortex carcinoma cell lines and certain lung adenocarcinoma cell lines display marked imbalances between the expression of lamins A and C (Paulin-Levasseur et al, 1988; Machiels et al, 1995). The fast growing small cell lung carcinomas also do not express A-type lamins whereas non-small cell lung carcinomas do (Broers and Ramaekers, 1994) reinforcing the suggestion of a relationship between expression of A-type lamins and growth rate and differentiation status.
In this study, we have used an extensive and unique panel of type-specific lamin antibodies to study lamin expression in human skin to look further at lamin expression in normal and diseased tissue. Our results reveal different expression patterns, not only between A-type and B-type lamins, but also between lamins A and C and between lamins B$_1$ and B$_2$ in different cell types. Furthermore, we report that while absence of lamin A is characteristic of basal cell skin carcinomas (BCCs), absence of lamin C occurs in only a sub-set of BCCs.

**MATERIALS AND METHODS**

**Antibodies**

A newly-generated antibody specific for lamin C was used in this study. This rabbit antiserum was raised against the last 8 amino acids of lamin C, including an N-terminal lysine as a linker (KHIVSGSRR). The peptide was coupled to keyhole limpet haemocyanin through primary amino groups using glutaraldehyde. The resulting protein-peptide conjugate was dialysed overnight at 4°C against PBS. The conjugate was then used to immunize a rabbit. Immune serum was screened by indirect immunofluorescence and then affinity purified against 10 mg of the lamin C peptide conjugated to CH Sepharose 4B as described in Harlow and Lane (1988).

All the primary antibodies to lamins used in this study are listed in Table 1, together with their sources and specificities. Rabbit antiserum to K667 was purchased from DAKO and used at a dilution of 1:50. All secondary antibodies were obtained commercially: FITC- and peroxidase-conjugated goat anti-mouse IgG, TRITC- and peroxidase-conjugated donkey anti-goat IgG and FITC-, TRITC- and peroxidase-conjugated sheep anti-rabbit IgG were all purchased from DAKO and used according to the manufacturer’s instructions. All dilutions were done in phosphate buffered saline (PBS) containing 1% newborn calf serum.

**Immunofluorescence**

HeLa epithelial cells or human dermal fibroblasts (HDF) were seeded at an initial density of 2 × 10$^4$ on 90 mm plates containing 13 mm diameter glass coverslips. The cells were grown to approximately 80% confluency in DMEM supplemented with 2 mM glutamine, 10 U ml$^{-1}$ penicillin, 100 μg ml$^{-1}$ streptomycin, plus 10% newborn (for HeLa cells) or fetal (for HDF cells) bovine serum. The cells were then fixed in methanol/acetone (1:1 v/v) for 10 minutes at 4°C and washed 3 times in PBS before immunostaining.

Biopsies of normal skin from different body sites (surgical trimmings from groin, dorsal-proximal finger, face and back areas) and of basal cell carcinomas (BCCs) from 16 individual patients were snap-frozen in liquid nitrogen. Frozen blocks were sectioned on a Reichert Jung cryomicrotome to give serial sections of approximately 6–7 μm thickness. Sections were collected on uncoated microscope slides and stored at −70°C. Prior to use they were removed from the freezer, thawed and dried rapidly in a stream of warm air. All staining procedures were then carried out at room temperature. The sections were incubated with 1% fetal calf serum in phosphate buffered saline (PBS) for 10 minutes to block non-specific protein binding, then rinsed in PBS for 5 minutes.

For immunofluorescence of cells or tissue sections, antibody incubations were all for 1 h at room temperature in a humidified environment, followed by 5 min wash in PBS. After the secondary antibody and wash, a final wash in water was used before mounting. Specimens were mounted in Mowiol containing 1 μg ml$^{-1}$ DAPI (4',6-diamidin-2-phenylindol-dihydrochloride) and 1 μg ml$^{-1}$ DABCO (1,4-diazabicyclo[2.2.2]octane). Immunofluorescence samples were examined with a Zeiss Axiosvert 10 microscope equipped for epifluorescence using ×63 N/A 1.4 and ×40 N/A 1.3 PlanNeofluar lenses. Images were collected with a 12 bit CCD camera using IPLab software.

**Immunoblotting for antibody characterization**

Three fragments of the lamin C gene encoding residues 1–171aa, 171–319aa and 319–572aa were cloned into pGEX expression vectors and expressed in E. coli strain BL21 and purified according to previously published protocols (Dyer et al, 1997; Pugh et al, 1997). Nuclear matrix preparations of HeLa cells were performed following previously published protocols of Mattern et al (1992).

Samples prepared for SDS-PAGE were resolved on a 12% polyacrylamide gel. Immunoblotting was carried out as described by Jenkins et al (1993) using the following primary antibodies (see Table 1): rabbit anti-lamin C (to lamin C) at 1:1000, JoL2 (to lamins A,C) used undiluted, JoL4 (to lamin A) used undiluted, LN43 (to lamin B$_1$) diluted to 1:1000 and affinity purified goat anti-lamin B$_1$ used at 1:500 (Santa Cruz). Detection of the primary antibody used antibodies conjugated to horseradish peroxidase as above. All antibody incubations were for 1 hour at room temperature. The blots were developed using the ECL detection method.

**RESULTS**

**Confirmation of antibody specificities**

The aim of this study was to distinguish between the expression and distribution of lamin subtypes and an essential first step was therefore to confirm the antibody specificities. On immunoblotting, the rabbit antiserum reacted strongly with fragments corresponding to peptide 319–572aa of lamin C but with no other fragment (Figure 1A). When tested against nuclear protein extracts from HeLa cells monoclonal antibodies specific for lamin A (JoL4 – Figure 1B, also 133A2) reacted with a single band migrating at 70 kDa. JoL2 against lamins A and C, reacted with two bands migrating at 70 kDa and 65 kDa respectively (Figure 1B, lane 1). The anti-peptide antiserum to lamin C reacted with a single band migrating at 67 kDa (Figure 1B, lane 4). The goat polyclonal antibody against lamin B$_2$ reacted with a single band migrating at 65 kDa (Figure 1B, lane 3). The goat polyclonal antibody against lamin B$_1$ reacted with a single band migrating slightly behind the lamin B band at 68 kDa (Figure 1B, lane 5). The expected mobilities of human lamins A, C, B$_1$ and B$_2$.
data confirm the lamin specificities indicated in Table 1 and especially that the rabbit polyclonal antibody generated for these studies specifically reacts with lamin C only.

**Distribution of lamins in normal human skin**

Immunofluorescence on frozen unfixed sections of skin from different body sites gave consistent results for each lamin. Typical examples (all from finger skin) are shown in Figure 3. Two different monoclonal antibodies against lamin A (JoL4 and 133A2) stained dermal fibroblasts and suprabasal epidermal cell nuclear envelopes intensely, but basal epidermal cells were mostly unstained (arrows Figure 3 D–E, Figure 4A). In contrast, JoL2 (anti-lamin A/C) stained dermal fibroblasts, suprabasal epidermal cells and the majority of basal epidermal cells. It was noticeable that a population of the basal epidermal cells were not stained by lamin A/C antibodies (Figure 3A–C; Figure 4C).

The goat anti-lamin B1 antibody stained suprabasal and basal epidermal cells uniformly. In contrast, dermal fibroblasts were mostly unstained (arrows, Figure 3G–I). The lamin B2 antibody LN43 stained all cell types in human skin (Figure 3J–L). The rabbit polyclonal antibody against lamin C (RtLC) gave rise to a staining pattern that was almost identical to JoL2 (anti-lamin A/C): dermal fibroblasts, suprabasal epidermal cells and a majority of basal epidermal cells were all stained. However, a small number of basal epidermal cells were unstained (arrows Figure 3 M–O).

Taken together, these data indicate differences in the expression of the lamin sub-types in different cell types within the epidermis. Lamin B1 is expressed uniformly in all nucleated cells within human skin (epidermis and dermis). Lamin B2 is expressed throughout the epidermis but as reported previously (Broers et al, 1997) was not detected in dermal fibroblasts. Lamin A appears to be expressed in dermal fibroblasts and in suprabasal epidermal layers, but most cells, in the basal layer did not express lamin A. Lamin C is also expressed in all dermal fibroblasts and in a majority of epidermal cells, but lamin C was still not detected in a small number of epidermal basal cells. Since JoL2 (A/C) and RtLC give identical staining patterns (this was confirmed by double immunofluorescence, not shown), but very different patterns from those observed with JoL4 and 133A2 (A), we infer that staining of basal epidermal cells with JoL2 indicates the presence of lamin C.

**Differences in the expression of lamin A and lamin C in epidermal basal cells**

In the epidermal basal cell layer, lamin A is not expressed in the majority of cells, whereas lamin C is absent from approximately 1 in every 40 cells. To investigate this phenomenon further, double indirect immunofluorescence was performed with monoclonal antibodies JoL2, JoL4, 133A2 and a rabbit polyclonal antibody against the proliferation marker Ki67. Whilst again 133A2 and JoL4 staining (lamin A) was absent from both Ki67-negative and Ki67-positive cells (Figure 4A, B) in the basal epidermis, JoL2 (A/C) staining was present in and apparently restricted to all Ki67-positive basal cells (arrowheads Figure 4C, D). These data suggest that A-type lamins are not expressed in a subpopulation of quiescent non-cycling basal cells in the epidermis, but that lamin C, but not lamin A, is up-regulated in cells that are stimulated to divide. Lamin A expression appears later in the process of differentiation.
Changes in the expression of lamins A and C in basal cell skin carcinomas

Changes in the expression of A-type lamins have been reported in a range of tumours. We investigated lamin expression in a collection of 16 basal cell skin carcinomas (BCC) taken from a variety of body sites including back, elbow and neck. In each instance lamin expression in normal (peri-tumoral) tissue and tumour lobes was compared. Typical results are displayed in Figures 6 and 7 while a summary of results is displayed in Table 2. All tumours retained expression of lamin B₁, and all except one expressed lamin B₂ also. The A-type lamins were more variable however. Most tumours expressed either lamin A (detected with JoL4 and 133A2) or lamin C but not both, although one tumour had neither. The A-type lamins were more variable however. Most tumours expressed either lamin A (detected with JoL4 and 133A2) or lamin C but not both, although one tumour had neither. The tumours could therefore be divided up into four groups on the basis of their lamin expression: A-negative (10/16 tumours), C-negative (5/16 tumours), A/C-negative (1/16 tumours) and A/B₂-negative (1/16 tumours). Finally, in one tumour from the A-negative group, lamin C was expressed but it was localized in the nucleolus rather than in the nuclear envelope (Figure 5; Table 2).

To determine whether lamin expression was correlated with rate of growth within the tumour, serial sections from the same tumours were also stained for Ki67 expression. Of the A-negative tumours, 9 out of 10 were highly proliferative as indicated by extensive Ki67 staining within the tumour lobe (Table 2). In

| Patient | Lamin A | Lamin A/C | Lamin C | Lamin B₁ | Lamin B₂ | Ki67 |
|---------|---------|-----------|---------|---------|---------|-----|
| 1.      | ++      | ++        | ++      | ++      | ++      | ++  |
| 2.      | ++      | ++        | ++      | ++      | ++      | ++  |
| 3.      | –       | +         | +       | ++      | ++      | ++  |
| 4.      | –       | +         | +       | ++      | ++      | ++  |
| 5.      | –       | +         | +       | ++      | ++      | ++  |
| 6.      | +/–     | +         | +       | ++      | ++      | ++  |
| 7.      | –       | +/–       | +       | ++      | ++      | ++  |
| 8.      | –       | +/–       | +       | ++      | ++      | ++  |
| 9.      | –       | +/–       | +       | ++      | ++      | ++  |
| 10.     | –       | –         | +       | ++      | ++      | ++  |
| 11.     | –       | s/–       | +       | ++      | ++      | s/– |
| 12.     | –       | –         | –       | ++      | ++      | ++  |
| 13.     | +       | –         | ++      | ++      | ++      | s/– |
| 14.     | +       | –         | ++      | ++      | ++      | s/– |
| 15.     | +       | –         |+++      | ++      | ++      | s/– |
| 16.     | ++      | ++        | +++     | ++      | ++      | +   |

Frozen sections of basal cell skin carcinomas were stained individually with type specific anti-lamin antibodies or with rabbit anti-Ki67. The level of expression of each antigen within tumour lobes was scored according to an arbitrary scale in which – indicates low level or absent staining while +++ indicates maximum staining. +/– indicates some positive some negative. Patient numbers are indicated in the left hand column. Nucleolar (lamin C patient 96/96) indicates that in this section anti-lamin C antibodies stained the nucleolus.
contrast, 4 out of the 5 C-negative tumours were slow growing as indicated by an absence of Ki67 staining within the tumour lobe (Table 2). Therefore as a general rule, absence of lamin A alone was correlated with rapid growth within the tumour, whereas absence of lamin C was correlated with slow growth within the tumour.

**DISCUSSION**

Until now there have been very few attempts to functionally discriminate between lamins A and C, largely because of the difficulty in doing so. The generation of an antibody specific to lamin C now allows a re-examination of the expression of A-type

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**Figure 3** Lamin expression in normal human finger skin. Frozen sections of normal human finger were stained with either JoL2 (anti-lamin A/C – panels A–C), 133A2 (anti-lamin A – panels D–F), goat anti-lamin B1 (panels G–I), LN43 (anti-lamin B2–panels J–K) or rabbit anti-lamin C (panels M–O). Sections were co-stained with DAPI to reveal nuclei. Panels A, D, G, J, M show DAPI stained images. Panels B, E, H, K and n show antibody stained images. Panels C, F, I, L and O show merged images. Arrows illustrate representative nuclei in each section that are unstained. Scale bars = 100 μm

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In this investigation, the absence of expression of one or other A-type lamin in the majority of a group of 16 basal cell carcinomas of the skin has been shown (the A-negative BCC type). Most tumours show a down-regulation of lamin A expression, while a smaller group of tumours (C-negative) gave negative results for lamin C expression. In one tumour neither A nor C lamins were detected. In another tumour, absence of lamin B2 expression accompanied absence of lamin A expression. In a third BCC, lamin C was present but abnormally located in the nucleolus.

Down regulation of A-type lamin expression is a feature of a number of tumour types including small cell lung carcinomas (Broers and Ramaekers, 1994), testicular cancer (Machiels et al, 1997) and Hodgkins disease (Jansen et al, 1997). In lung cancer, changes in the level of lamin expression (Broers et al, 1993) have been linked to expression of v-rasH (Kaufman et al, 1991), but there is apparently no common mechanism by which down regulation of lamin expression occurs in tumours. The lamin A gene has been mapped to 1q12.1-q23, a region of chromosome 1 that is commonly rearranged in tumours (Kamat et al, 1993). Gene deletion (Kamat et al, 1993) as well as transcriptional and

Figure 4 Lamin A and C expression in basal epidermis. Frozen sections of finger skin were co-stained with JoL2 (anti-lamin A/C), Ki67 and DAPI (panels A and B) or JoL4 (anti-lamin A), Ki67 and DAPI (panels C and D). Each panel represents a two colour merged image in which the anti-lamin antibody stain (in green) is superimposed over the DAPI stain (in blue – panels A and C) or the Ki67 stain (in red – panels B and D). Arrows indicate cells that are Ki67 negative: arrowheads indicate cells that are Ki67 positive. Scale bars = 100 μm

Figure 5 Lamin expression in basal cell skin carcinomas. Frozen sections from BCCs were co-stained with one of JoL 4 (anti-lamin A – panels A and B), JoL 2 (anti-lamins A/C – panels C and D), rabbit anti-lamin C (panels E and F), goat anti-lamin B1 (panels G and H) or LN43 (anti-lamin B2 – panels I and J) and DAPI. In each panel the antibody stain (in red or green) is superimposed over the DAPI image (in blue). The left hand panels show staining of patient number 46/95 that is a representative lamin A negative, lamin C positive tumour. The right hand panels show an atypical lamin A negative, lamin C positive tumour (number 96/96) in which lamin C is localized in the nucleolus (arrows panel F). Scale bars = 100 μm
post-transcriptional mechanisms (Lebel et al., 1987; Machiels et al., 1996) have all been implicated in the loss of lamin A or C expression. Therefore an important question is whether changes in lamin expression reflect de-differentiation within the tumour or is a requisite of tumour progression.

10 out of the 16 basal cell carcinoma tumour samples included in this investigation did not show lamin A expression. Lamin A was not detected in most epidermal basal cells in normal tissue, suggesting that these cells could be the originators of the A-negative tumours. Thus the lamin A detected in 6 samples could reflect up-regulation in these tumours. Finally, 5 samples showed an absence of lamin C in tumour lobes, although lamin C was expressed in normal proliferating basal cells of the epidermis, suggesting that this could reflect downregulation of lamin C expression. These data therefore suggest a complex pattern of lamin A/C expression that is not controlled by a single mechanism.

In the majority of cases, absence of lamin A might be explained by the origin of the tumour (i.e. from lamin-A negative basal epidermal cells). However previous studies have shown that expression of lamin A is down-regulated as mouse fibroblasts progress from a quiescent to a proliferating state (Pugh et al., 1997). The majority of tumours failing to stain with lamin A antibodies were hyperproliferative, as seen by Ki67 staining, and an alternative explanation for the absence of lamin A in these tumours is through cell-cycle dependent mechanisms. Tumours displaying absence of lamin C were generally slower growing (less Ki67). Changes in lamin C expression resulting from changes in cell cycle status have not been detected in earlier studies (Dyer et al., 1997; Pugh et al., 1997). Therefore, the absence of lamin C in slow-growing tumours was somewhat surprising but again indicates underlying selective changes in lamin expression associated with the rate of growth in the tumour. Since lamin A and lamin C are alternatively spliced variants of the same gene (Lin and Worman, 1993), the differences in expression of these two lamins must arise through post-transcriptional mechanisms (Lanoix et al., 1992). The availability of specific antibodies against lamin C allows for further study of this phenomenon which will be essential in order to understand why the expression of these two lamins is linked to the rate of tumour growth.

Lamin A has been reported as an in vitro binding partner for the tumour suppressor protein p110RB (Ozaki et al., 1994), and p110RB is tightly associated with nuclear substructures in its hypophosphorylated form (Templeton et al., 1991; Mancini et al., 1997). Although lamins are normally distributed at the nuclear envelope, more recent evidence has shown that lamin A is also located at a number of intranuclear sites (Goldman et al., 1992; Bridger et al., 1993), which could provide nuclear anchorage sites for p110RB. If as suggested, nuclear anchorage is important for the function of p110RB in transcriptional silencing (Mittnacht and Weinberg, 1991), an absence of lamin A may favour unregulated cell division. Therefore, its presence or absence may directly influence the proliferative status of a tumour. That lamin A is absent in hyperproliferative basal cell skin carcinomas supports this hypothesis.

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