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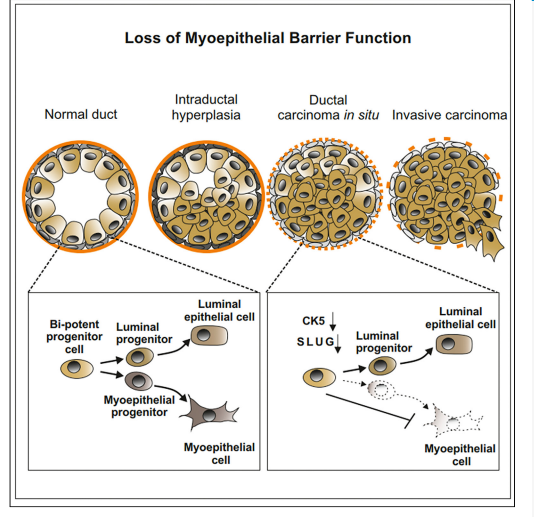
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# Article

# Cytokeratin 5 determines maturation of the mammary myoepithelium



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#### Highlights

Cytokeratin 5 (CK5) impacts mammary epithelial cell lineage differentiation

CK5 loss at pre-invasive stage causes impaired maturation of myoepithelial cells

CK5 loss causes SLUG downregulation and differentiation bias in mammary progenitors

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# Article

# Cytokeratin 5 determines maturation of the mammary myoepithelium

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#### SUMMARY

At invasion, transformed mammary epithelial cells expand into the stroma through a disrupted myoepithelial (ME) cell layer and basement membrane (BM). The intact ME cell layer has thus been suggested to act as a barrier against invasion. Here, we investigate the mechanisms behind the disruption of ME cell layer. We show that the expression of basal/ME proteins CK5, CK14, and  $\alpha$ -SMA altered along increasing grade of malignancy, and their loss affected the maintenance of organotypic 3D mammary architecture. Furthermore, our data suggests that loss of CK5 prior to invasive stage causes decreased levels of Zinc finger protein SNAI2 (SLUG), a key regulator of the mammary epithelial cell lineage determination. Consequently, a differentiation bias toward luminal epithelial cell type was detected with loss of mature,  $\alpha$ -SMA-expressing ME cells and reduced deposition of basement membrane protein laminin-5. Therefore, our data discloses the central role of CK5 in mammary epithelial differentiation and maintenance of normal ME layer.

#### INTRODUCTION

Mammary gland parenchyme forms a ducto-lobular tree with a bilayered epithelium. The inner layer of luminal epithelial (LE) cells is surrounded by a basal/myoepithelial layer, which comprises contractile myoepithelial (ME) cells, mammary stem cells, and epithelial progenitor cells, delimited by the basement membrane (BM) from the connective tissue stroma. The basal cell layer is able to regenerate the whole mammary gland epithelial tree (Böcker et al., 2002; Boecker and Buerger, 2003; Van Keymeulen et al., 2011). Development of LE and ME cells occurs in a complex hierarchical manner from the basal progenitors that can differentiate into both epithelial cell types depending on numerous signaling pathways and hormonal stimuli (Arendt and Kuperwasser, 2015; Böcker et al., 2002; Boecker and Buerger, 2003; Boecker et al., 2018; Van Keymeulen et al., 2011). This epithelial differentiation process can be followed as changes in cell-type specific protein expression patterns, including expression of distinct cytokeratin (CK) family members (Böcker et al., 2002; Boecker and Buerger, 2003; Boecker et al., 2018). Less than 5% of the mammary basal cells represent mammary stem cells that express CK5 without luminal epithelial (LE) markers CK8/18/19 or myoepithelial (ME) marker α-SMA (Böcker et al., 2002; Fu et al., 2020). Expression of CK5 is also detected in the progenitor cells that can differentiate into mature luminal or ME cells, lacking the expression of CK5 (Böcker et al., 2002; Boecker and Buerger, 2003; Boecker et al., 2018). Additionally, progenitor cell activity has been attributed to cells expressing CK14 simultaneous with LE or ME markers (Arendt et al., 2014; Boecker et al., 2018; Fridriksdottir et al., 2017; Villadsen et al., 2007). While CKs are filament-forming proteins that mechanically support the cell structure, they have also been attributed to other regulatory functions, such as coordination of nuclear morphology, cell proliferation and apoptosis (Bozza et al., 2018; Iver et al., 2013; Pan et al., 2013; Weng et al., 2012). Whether basal CK5 has other than structural roles in mammary stem and progenitor cells has not been assessed.

Stem or progenitor cells have been suggested to act as targets for neoplasia initiating transformation (Jiang et al., 2010; Molyneux et al., 2010; Reya et al., 2001). Most mammary carcinomas represent malignant intraductal hyperplasia of epithelial cell origin. Non-invasive and invasive intraductal proliferative lesions are distinguished. Non-invasive lesions comprise usual ductal hyperplasia (UDH), atypical ductal hyperplasia (ADH) and ductal carcinoma *in situ* (DCIS) (Schnitt et al., 2012). Long-term follow-up studies have shown

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that in time DCIS cases without treatment may develop into invasive carcinomas and may eventually lead to distant metastasis. The percentage of transformed cases varies amongst others according to follow-up time and grade of lesion, with values for low-grade ranging between 18 and 50%, for intermediate grade between 32 and 33% and for high-grade between 17.6 and 67%, respectively (Collins et al., 2005; Maxwell et al., 2018; Ryser et al., 2019; Sanders et al., 2005, 2015). The invasive stage is determined when the boundary provided by ME cells and BM is disrupted. Therefore the ME cell layer has been suggested to act as a barrier against invasion.

During the neoplastic transformation, mammary epithelial cells undergo alterations in their gene expression patterns (Allinen et al., 2004). Unfortunately, it has not been possible to identify distinct markers to predict this transformation from in situ to invasive disease (Yeong et al., 2017). DCIS-associated ME cells display immunophenotypic differences in comparison to ME cells surrounding normal structures. Several markers, such as basal CKs, α-SMA, SMMHC, calponin, p63, p75, maspin, WT-1, and CD10 have been demonstrated to decrease prior to the invasive stage (Chocteau et al., 2019; Guelstein et al., 1993; Hilson et al., 2009; Kalof et al., 2004; Rohilla et al., 2015; Werling et al., 2003; Wetzels et al., 1989; Zhang et al., 2003). A sequential disappearance has been shown for p63, calponin, and  $\alpha$ -SMA. The loss of  $\alpha$ -SMA expression is linked to later events, and taking place just before invasion (Russell et al., 2015). Recently, loss of  $\alpha$ -SMA was also shown to compromise the barrier made by ME cells (Sirka et al., 2018), suggesting that the ME layer acts as a mechanical barrier and that the contractile potential, mediated by  $\alpha$ -SMA, is important for its protective function. Besides displaying physical hindrance, ME cells are known to participate in the production of BM and regulation of matrix metalloproteinases, further supporting their importance against invasion (Gudjonsson et al., 2002; Jones et al., 2003; Sánchez-Céspedes et al., 2016; Sarper et al., 2017). Alterations in DCIS-associated ME cells have been demonstrated by gene expression profiling (Allinen et al., 2004). However, the mechanisms behind the disruption of the ME layer during malignant progression are not well understood.

To better understand the role of an intact ME cell layer and mechanisms behind its maintenance, we have utilized a comparative canine mammary tumor model for human breast carcinomas. Canine and human mammary tumors share similarities in their epidemiology, etiology, histomorphology, biological behavior, and molecular biology. In addition, most mammary carcinomas in humans and canines represent malignant hyperplasias of ductal epithelial cells (Goldschmidt et al., 2011; Klopfleisch et al., 2011; Rasotto et al., 2014; Rivera and von Euler, 2011; Sorenmo et al., 2011; Uva et al., 2009). Using immunohistochemistry, we compared expression patterns of basal/ME markers in untransformed canine mammary tissue sections with non-invasive intraductal epithelial proliferative lesions of UDH, ADH, and DCIS. We observed that in the basal/ME cells the expression of cytoskeletal proteins  $\alpha$ -SMA, CK5, and CK14 slightly responded to intraductal proliferations according to the ductal segment and type of proliferative lesion. At the invasive stage, the expression of these specific markers was absent, coinciding with the disruption of the intact ME cell layer. Furthermore, our cell biological experiments with primary canine mammary epithelial cells and the human mammary epithelial cell line showed that the loss of CK5, and to a lesser extent CK14, from the basal progenitor population affected maturation of the progenitors into functional, contractile ME cells. Simultaneously, a differentiation bias toward the luminal epithelial cell type was detected with loss of normal 3D mammosphere morphology and reduction in the basement membrane protein laminin-5. Importantly, loss of CK5 was associated with downregulation of transcriptional repressor Zinc finger protein SNAI2 (SLUG), an important regulator of the mammary epithelial cell lineage determination. In conclusion, our data suggest that CK5 impacts lineage specific differentiation and in this way may direct the formation of a normal ME layer, subsequently affecting the maintenance of a normal BM layer and mammary organostructural homeostasis. Hence, our findings expand our understanding of the carcinogenetic mechanisms at the pre-invasive stage and the development of phenotypic heterogeneity in mammary carcinomas.

#### RESULTS

# Basal/myoepithelial markers CK5, CK14 and α-SMA display specific expression patterns according to ductal segment and type of intraductal hyperplasia

In women and female canines, invasive mammary carcinomas have been proposed to originate from the terminal duct lobular unit (TDLU). The TDLU comprises a lobule with acini (terminal ductules) and intralobular terminal duct together with an extralobular terminal duct, which drains into a larger interlobular duct (Figure S1A). In these structures, ME cells display spatial differences in their morphology and immunophenotype, and molecular alterations have been demonstrated in DCIS-associated ME cells (Allinen et al.,

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2004; Chocteau et al., 2019; Hilson et al., 2009; Rønnov-Jessen et al., 1996; Russell et al., 2015; Sánchez-Céspedes et al., 2016).

To establish these observations quantitatively, we first determined the 3.3'-diaminobenzidine-tetrahydrochloride (DAB) chromogen staining intensity (DABi) values for the basal/myoepithelial markers CK5, CK14 and  $\alpha$ -SMA from normal intralobular terminal ductal segments and extralobular terminal/interlobular ductal segments using scanned immunohistochemical canine mammary tissue serial sections (Figures S3A and S3B). Slight but statistically significant difference was observed between the ductal segments for  $\alpha$ -SMA, with normal intralobular terminal ducts showing lower DABi values in comparison to extralobular terminal/interlobular ductal segments. However, the DABi values for CK5 and CK14 were not statistically significant between different segments (Figure S1B). Hence, our data demonstrate quantitatively that the expression pattern of basal/ME marker  $\alpha$ -SMA differs in a spatial manner in the normal ductal segments.

 $\alpha$ -SMA, CK5, and CK14 are known to be lost from the basal/ME layer prior to the invasive stage (see e.g. Rohilla et al., 2015; Russell et al., 2015; See also Figure 1A for  $\alpha$ -SMA example). To evaluate how the expression of these proteins is altered with an increasing grade of malignancy, we explored the DABi of these markers in non-invasive intraductal proliferative lesions of UDH, ADH and various grades of DCIS, and compared them to one another and to the normal ductal segments in the same canine patients (Figures 1B, 1C, S2A, and S2B). This analysis showed that in the extralobular terminal/interlobular ducts, statistically significant difference can be determined only for CK5 with I-G DCIS values slightly lower than in the corresponding normal ducts (Figure 1C).

In the intralobular terminal ductal segments CK5 exhibited lower values in florid UDH compared to normal and higher values in I-G DCIS compared to florid UDH. Lower values were determined for α-SMA in normal compared to I-G DCIS, and in mild-to-moderate UDH compared to I-G DCIS. CK14 values did not show statistically significant differences (Figure S2B). Taken together, these data show slight spatial differences in the expression of ME marker α-SMA between the normal ductal segments. The other comparative results suggest that cytoskeletal CK5 and α-SMA in the basal/ME layer may respond to non-invasive intraductal proliferative lesions in a spatial- and lesion-type-dependent manner.

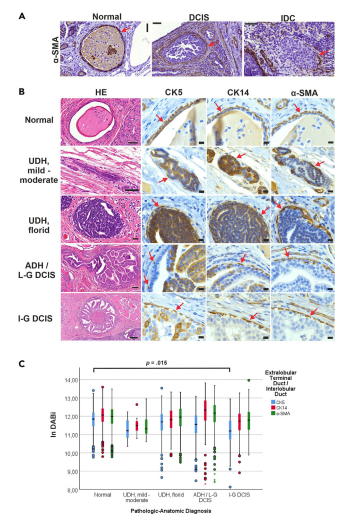
#### Loss of α-SMA, CK5 and CK14 leads to abnormal 3D mammosphere formation

In both humans and canines, CKs 5 and 14 as well as  $\alpha$ -SMA are expressed in the mammary basal/ME layer, but lost upon invasion (see e.g. Rohilla et al., 2015; Russell et al., 2015; Chocteau et al., 2019; Figure 1A). To understand how the loss of these specific markers would contribute to the properties of the ME layer and to the overall morphology of mammary epithelial structures, we isolated CD24<sup>+</sup> -epithelial cells with fluorescence-activated cell sorting (FACS) from primary canine mammary organoids (Figures S3C-S3F). The isolated population comprised of basal and luminal cells (Figure S3G) (Sleeman et al., 2006). These primary epithelial cells were targeted with CK5, CK14, or α-SMA-specific siRNAs in 3D Matrigel cultures (Figures 2A and S4A). Loss of each of these proteins led to significantly altered mammosphere morphology with a larger diameter (Figures 2A and 2B). Furthermore, we depleted these proteins from human MCF10A mammary epithelial cells, containing both basal and luminal cell types (Krause et al., 2008; Bhat-Nakshatri et al., 2010; Sarrio et al., 2012; Liu et al., 2014; Sokol et al., 2015; Qu et al., 2015; Miller et al., 2018). As these cells in long-term 3D cultures express markers against both luminal and basal/ME cells and produce laminin-5 to the forming basement membrane (Figures 2C and S4B; See also refs. Debnath et al., 2003; Gaiko-Shcherbak et al., 2015; Pinto et al., 2011; Qu et al., 2015; Pseftogas et al., 2020), we found this cell line as a useful model to target basal/ME proteins. Similar to canine mammary epithelial cells, loss of CK5, CK14, or α-SMA from human mammary epithelial cell cultures by specific siRNAs led to significantly enlarged mammosphere structures of abnormal morphology (Figures 2D and 2E).

To further explore whether the loss of these specific CKs could affect the morphology of 3D structures by impacting the epithelial differentiation process, we isolated the CD49f<sup>+</sup> EpCAM<sup>-</sup> population (Eirew et al., 2008; Stingl et al., 2001), enriched for basal progenitors, from the MCF10A cell line (Figures S4C–S4E). These progenitor cells were targeted by lentiviral-based RNA interference to knock down CK5 and CK14 (Figures S4F, S4G, and S5A). Similar to siRNA experiments these knock down (KD) cells in a 3D environment formed larger mammospheres with abnormal morphology (Figures 2F,S5B, and S5C). These results







# Figure 1. Intraductal hyperplasia in the extralobular terminal duct/interlobular ductal segment is associated with modest basal myoepithelial response that is lost prior to invasive stage

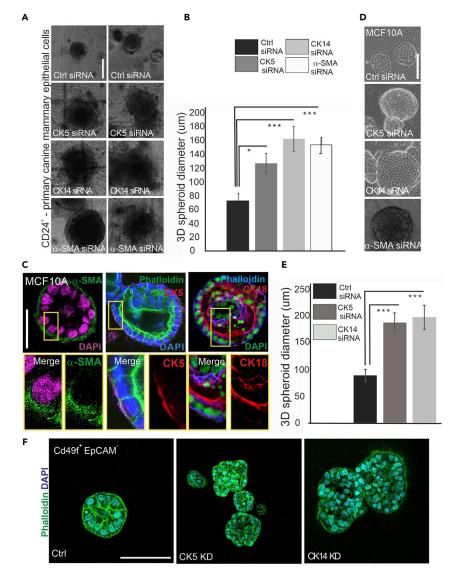
(A) Loss of myoepithelial cell layer prior to invasive stage. Canine mammary myoepithelial layer visualized with immunohistochemical staining for α-SMA (red arrows) in normal interlobular duct (left), ductal carcinoma *in situ* (DCIS; middle) and invasive ductal carcinoma (IDC; right). Bar 50 μm.

(B) Consecutive canine mammary FFPE tissue sections of normal, mild-to-moderate usual ductal hyperplasia (UDH, mildmoderate), florid usual ductal hyperplasia (UDH, florid), atypical ductal hyperplasia (ADH), low-grade ductal carcinoma *in situ* (L-G DCIS) and intermediate-grade ductal carcinoma *in situ* (I-G DCIS) were stained using hematoxylin-eosin (HE, far left) and the basal myoepithelial markers CK5 (middle left), CK14 (middle right) and α-SMA (far right). Representative images of the lesions are shown. Red arrows indicate basal myoepithelial layer as distinct from intraluminal cellular hyperplasia. Bar for HE in normal and mild-to-moderate UDH 50 µm, florid UDH 20 µm, ADH/L-G DCIS 50 µm, I-G DCIS 100 µm and in all IHC 10 µm.

(C) Boxplot of the extralobular terminal / interlobular ductal cellular In-transformed DAB chromogen staining intensity (DABi) values for normal CK5 n(cells) = 1203, CK14 n = 1469,  $\alpha$ -SMA n = 1765; mild-to-moderate UDH CK5 n(cells) = 12, CK14 n = 12,  $\alpha$ -SMA n = 16; florid UDH CK5 n(cells) = 248, CK14 n = 403,  $\alpha$ -SMA n = 304; ADH/L-G DCIS CK5 n(cells) = 268, CK14 n = 383,  $\alpha$ -SMA n = 383; I-G DCIS CK5 n(cells) = 203, CK14 n = 258,  $\alpha$ -SMA n = 319. Canine patient n = 7. Black middle line within box represents median. Height of box is interquartile range (IQR), representing 75<sup>th</sup> and 25<sup>th</sup> percentiles, respectively. Whiskers represent the lowest and highest data within the 1.5 x IQR of the lower and upper quartiles, respectively. Circles represent outliers. Linear mixed model with random intercepts for canine individual and ductal segment was used. Pairwise comparison over all the individual companions was implemented with Bonferroni's multiple comparisons correction. The level of significance was defined as p < 0.05. Only statistically significant mean differences are indicated. CK5 expression was significantly decreased in I-G DCIS compared to normal (p = 0.015). See also Figures S1–S3.

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#### Figure 2. Loss of CK5, CK14 or α-SMA affects the homeostasis of mammary epithelial structures

(A) Depletion of primary canine mammary epithelial cells (CD24<sup>+</sup>) by specific siRNAs against CK5, CK14, and  $\alpha$ -SMA in 3D Matrigel. Two panels of representative bright field images after 2 weeks of culture are shown. Bar 100  $\mu$ m.

(B) Quantification of canine 3D mammary organoid diameters from ctrl, CK5-, CK14-, and SMA-depleted samples, related to Figure 3A. Mean ( $\pm$ SEM) is shown; n(ctrl) = 31, n(CK5 siRNA) = 51, n(CK14 siRNA) = 31, n( $\alpha$ -SMA siRNA) = 33 \*\*\*P<0.001 (Mann–Whitney–Wilcoxon rank-sum test).

(C) 3D-structures of MCF10A cells in Matrigel display both luminal and basal markers. Human mammary epithelial cells were grown for two weeks in Matrigel, fixed with PFA, and stained with specific antibodies against  $\alpha$ -SMA, CK5 and CK18. Phalloidin was used to visualize actin cytoskeleton and DAPI for nuclei. Bar 50  $\mu$ m.

(D) MCF10A human mammary epithelial cells were depleted for CK5, CK14, and α-SMA by specific siRNAs in 3D Matrigel cultures. Samples were grown for two weeks and fixed with PFA for analyses. CK5- and CK14-depleted mammosphere samples were clearly larger than ctrl 3D structures. Bar 100 μm.

(E) Quantification of the 3D mammosphere diameter from ctrl, CK5- and CK14-depleted samples. Mean ( $\pm$ SEM) is shown; n(ctrl) = 15, n(CK5 siRNA) = 15, n(CK14 siRNA) = 15; \*\*\*P<0.001 (Mann–Whitney–Wilcoxon rank-sum test).

(F) Depletion of CK5 and CK14 from 3D cultures CD49f<sup>+</sup> EpCAM<sup>-</sup>/basal progenitor-enriched MCF10A mammary epithelial cell population. Cultures were maintained for two weeks, after which they were fixed with PFA and stained with Phalloidin (green) and DAPI (blue). Representative immunofluorescence images of the 3D mammospheres are shown. Bar 100 μm. See also Figures S3–S5.





indicate that basal/ME proteins CK5, CK14, and  $\alpha$ -SMA are important for the maintenance of normal mammary organomorphology, at least in the utilized *in vitro* 3D models.

#### Cytokeratin 5 determines maturation of the myoepithelial cells

CKs play a role in the mechanical resistance of epithelial cells (Sanghvi-Shah and Weber, 2017). In line with that, we observed a decrease in the elastic modulus of CK5 KD and CK14 KD cells in comparison to control (CD49f+ EpCAM-/basal progenitor-enriched MCF10A cells) in indentation experiments using an atomic force microscope (AFM). The elastic modulus histograms could be fitted with three Gaussian distributions, revealing a different mechanical behavior at distinct indentation spots within the same cell type (Figures 3A–3D). The elastic modulus values at the peaks of each distribution were significantly lower for CK5 KD and CK14 KD cells when compared with control cells (Figure 3D).

While CK5 and CK14 clearly maintain mechanical properties of the mammary basal/ME layer, it has not been assessed whether they could have additional regulatory roles in the progenitor cells. To understand the role of these CKs in the regulation of mammary progenitor cells, we utilized CD49f<sup>+</sup> EpCAM<sup>-</sup>/basal progenitor-enriched MCF10A cells that were depleted for CK5 and CK14, and analyzed the expression of several markers by Western blotting (WB) and by immunofluorescence (IF) stainings (Figures 3E-3G and S6A-S6C). The WB results showed that CK5 KD cells, and to a lesser extent CK14 KD cells, displayed decreased levels of ME cell marker α-SMA, while the luminal epithelial marker CK18 was slightly increased in CK5 KD cells and the luminal epithelial marker CK19 decreased (Figures 3E-3G and S6A-S6C). Additionally, CK5 KD cells displayed lower levels of ME cell markers vimentin, smooth muscle myosin heavy chain (SMMHC) and calponin 1 (Figures S6B and S6C). Similar results showing downregulation of vimentin and α-SMA were obtained by utilizing specific siRNAs against CK5 in MCF10A cell line (Figures 3H and 3I). In addition, we performed combined CK5/CK14 siRNA experiments, but could not see higher depletion of a-SMA upon the combined siRNA treatment in comparison to CK5 depletion alone (Figures S6D and S6E). Interestingly, depletion of α-SMA seemed to reciprocally downregulate CK5 and vimentin, indicating a feedback loop mechanism in between these proteins (Figures S6F and S6G). These results suggest that CK5 has a major role in the maturation process of ME cells, the loss of CK5 leading to a differentiation bias toward the CK18<sup>+</sup> luminal epithelial cell type. As CK5 and CK14 are known to heterodimerize, it may also be possible that the milder impact of CK14 depletion goes through CK5.

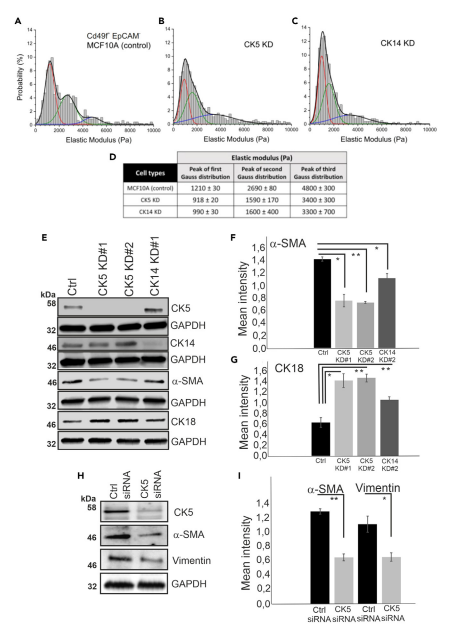
# Loss of Cytokeratin 5 impairs junctional integrity and affects deposition of basement membrane proteins

CKs are linked to integrin- and cadherin-based adhesions, and have been associated with regulation of these cell adhesive structures (Sanghvi-Shah and Weber, 2017). As CK5, and to a lesser extent CK14, were found to affect the maturation of ME cells, we wanted to assess whether loss of these proteins could also impact the resistance of the ME layer through cell adhesive structures. The levels of the ME-specific cell-cell contact proteins Dsg3 and P-cadherin (Daniel et al., 1995; Runswick et al., 2001) were determined from lysates of both ctrl (CD49f<sup>+</sup> EpCAM<sup>-</sup>/basal progenitor-enriched MCF10A cells) and the corresponding CK5 and CK14 KD cell lines (Figures 4A and 4B). Both markers were significantly decreased upon loss of CK5, while loss of CK14 did not seem to play a role in maintaining their levels (Figures 4A and 4B). More detailed immunofluorescence analyses of Dsg3-stained fully confluent epithelial monolayers revealed that Dsg3 partially lost its junctional pattern in CK5-deficient cells and that the cell-cell junctions appeared less mature, with spiky protrusions (Figures 4C and 4D). In a 3D environment, CK5 KD cells displayed lower Dsg3-staining pattern, while CK14 KD cells had many randomly localized cells with junctional Dsg3 within the morphologically abnormal 3D structures (Figure S7A). Staining of luminal marker E-cadherin from 3D mammospheres was, however, prominent in all samples but the distribution of E-cadherin positive cells in CK5 and CK14 KD spheroids was clustered and abnormal in comparison to the ctrl 3D mammospheres (Figures 4E and 4F). As with CK5 loss, slight decrease in Dsg3 and P-cadherin levels was detected upon  $\alpha$ -SMA-depletion by siRNA in MCF10A cells (Figure 4G). These data indicate that CK5 may play a role in the integrity of ME cell junctions at least through P-cadherin and Dsg3, and that bidirectional signaling within the basal/ME layer may be important for the overall maintenance of the epithelial cell populations.

As CK5 loss affected cell-cell adhesions, we further studied whether its downregulation would play a role in the regulation of cell-substrate adhesions. Immunofluorescence stainings with vinculin antibody in ctrl (CD49f<sup>+</sup> EpCAM<sup>-</sup>/basal progenitor-enriched MCF10A cells) or CK5-deficient cells showed slightly more prominent vinculin-based cell-substrate adhesions, while vinculin at the cell-cell contacts was showing







#### Figure 3. Loss of CK5 affects maturation of myoepithelial cells

(A–C) Elastic moduli of cells determined in cell indentation experiments by AFM. Elastic modulus histograms for (A) Control (CD49f<sup>+</sup> EpCAM<sup>-</sup> -enriched basal progenitors from MCF10A cells), (B) CK5 KD cells and (C) CK14 KD cells. The histograms were fit with 3 Gaussian distributions (black lines). Each Gaussian distribution is shown separately (red, green, and blue lines).

(D) Elastic modulus values for the peaks of the Gaussian distributions of Control (CD49 $f^+$  EpCAM $^-$  -enriched basal progenitors from MCF10A cells), CK5 KD, and CK14 KD cells.

(E) Western blot analyses on cell lysates from ctrl and CK5-and CK14-depleted cell lines showed downregulation of α-SMA, as detected by specific antibody. In contrast, luminal marker CK18 was slightly elevated in the corresponding cell lysate samples.

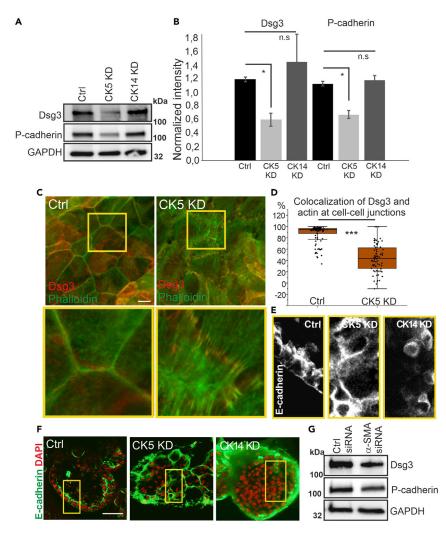
(F and G) Quantification of the  $\alpha$ -SMA and CK18 Western blot experiments. Mean ( $\pm$ SEM) is shown; n = 3; \*P<0.05, \*\*P<0.01 and \*\*\*P<0.001 (paired ttest).

(H) Ctrl siRNA-treated MCF10A cells and MCF10A cells depleted for CK5 siRNA were analyzed in Western blotting by specific antibodies against  $\alpha$ -SMA and vimentin. GAPDH was used as a loading control.

(I) Quantifications of the Western blots, related to Figure 3H. Mean ( $\pm$ SEM) is shown; n = 3; \*P<0.05 and \*\*P<0.01 (paired ttest). See also Figure S6.







#### Figure 4. Loss of CK5 affects cell-adhesive structures

(A) Western blot analyses on cell lysates from ctrl and CK5-and CK14-depleted cell lines showed downregulation of Dsg3 and P-cadherin upon loss of CK5 but not CK14. GAPDH is used as a loading control.

(B) Quantification of the Dsg3 and P-cadherin Western blot experiments, related to Figure 4A. Mean ( $\pm$ SEM) is shown. n = 4 \*P<0.05 (paired ttest); n.s.= not significant.

(C) Ctrl (CD49f<sup>+</sup> EpCAM<sup>-</sup>, enriched for basal progenitors) and CK5 KD cells were used in immunofluorescence microscopy of fully-confluent monolayer cultures. Specific antibody against Dsg3 was used. Actin cytoskeleton was visualized with Phalloidin and nuclei were stained with DAPI. Magnifications of the cell-cell junction areas, indicated with yellow boxes, are shown below. Bar 20  $\mu$ m.

(D) Quantification of the colocalization in between Dsg3 and actin at cell-cell junctions. n(ctrl) = 91, n(CK5 KD) = 75. The amount of colocalization (%) is shown as box plot with inner and outlier points and mean. \*\*\*P<0.001 (paired ttest). (E and F) (E) CD49f<sup>+</sup> EpCAM<sup>-</sup>/basal progenitor-enriched MCF10A mammary epithelial cells, CK5 KD and CK14 KD cells were culture in 3D Matrigel for two weeks, after which they were fixed with PFA and stained with E-cadherin. Nuclei were visualized with DAPI. Magnifications of E-cadherin stainings in gray scale are shown in panel (E) and full images with the indicated magnified areas (yellow boxes) are shown below in panel (F). Bar 25 um.

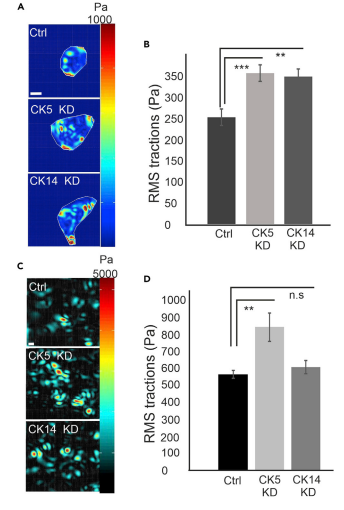
(G) Depletion of  $\alpha$ -SMA was performed with specific siRNAs in MCF10A cultures for four days. Cellular lysates from control siRNA and  $\alpha$ -SMA siRNA treated cells were used in Western blotting and a specific antibody against Dsg3 was utilized. GAPDH was used as a loading control. See also Figure S7.

punctate, immature type adhesive structures (Figure S7B). As vinculin has an established role in mechanotransduction (Goldmann, 2016), we tested whether cell-exerted forces would be altered in the CK5 KD progenitor cell lines. Traction force imaging experiments showed an increase in the actomyosin-mediated cellsubstrate forces in both single cells and monolayers but no changes in the cell doublets (Figures 5, S7C, and

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#### Figure 5. Loss of CK5 and CK14 impact cellular force production

(A) Traction force microscopy with ctrl, CK5 KD and CK14 KD cell lines showed altered cell-substrate forces upon CK5 and CK14 depletion. Representative force maps of ctrl, CK5 and CK14 knock-down cells are shown. Bar 20  $\mu$ m. (B) Quantification of the traction force microscopy experiments, related to Figure 5A, showed elevated cell-substrate forces upon loss of CK5 and CK14. Mean ( $\pm$ SEM) is shown. n(ctrl) = 32, n(CK14 KD) = 40, n(CK5 KD) = 33; \*P<0.05 (Mann-Whitney-Wilcoxon rank-sum test).

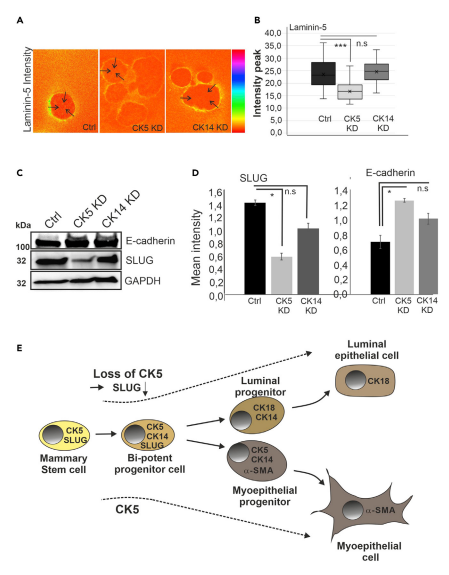
(C) Representative examples of monolayer force microscopy maps of ctrl, CK5 KD and CK14 KD cell sheets. Bar 40  $\mu$ m. (D) Quantification of the monolayer force microscopy experiments, related to Figure 5C, showed elevated cell-substrate forces upon loss of CK5. Mean ( $\pm$ SEM) is shown. n(ctrl) = 19, n(CK5 KD) = 16, n(CK14 KD) = 16; \*\*P<0.01; n.s= not significant (Mann–Whitney–Wilcoxon rank-sum test). See also Figure S7.

S7D). These results indicate that the lack of CK5 and CK14 may be counteracted by the redistribution of intercellular forces, and that loss of these cytokeratins may lead to redistribution of cellular forces.

Finally, to reveal whether CK5 could impact the barrier against transformed luminal cells also through basement membrane formation, we stained 3D mammosphere cultures with laminin-5. Fully mature MCF10A acinar structures in 3D are known to produce laminin-5 to the basement membrane (Gaiko-Shcherbak et al., 2015). While we detected this layer both in MCF10A and Cd49f+ EpCAM-cultures (Figures S4B and 6A), CK5 deficient cultures were displaying significantly decreased amounts of laminin-5 around the spheroids as visualized by the intensity maps of laminin-5-stainings (Figures 6A,6B, andS8A). However, CK14 KD cultures did not alter significantly from the ctrl cultures. Additionally, decreased levels of laminin-5 were detected in Western blot experiments, performed from CK5 siRNA-treated MCF10A cells (Figures S8B and S8C). These results act as additional proof for the observations that loss of CK5 not only







#### Figure 6. Depletion of CK5 affects laminin-5 production and Zinc finger protein SNAI2 (SLUG2) levels

(A) CD49f<sup>+</sup> EpCAM<sup>-</sup>/basal progenitor-enriched MCF10A mammary epithelial cells, CK5 KD and CK14 KD cells were culture in 3D Matrigel for two weeks, after which they were fixed with PFA and stained with laminin-5 (See also Figure S8A). Intensity maps were created in Fiji. Lineprofiles were drawn from the edge of the spheroid toward the center. 3–5 lineprofiles were drawn on each spheroid for the analyses of laminin-5 intensity.

(B) Peak values on point 3 from lineprofiles were utilized for further analyses. Values for ctrl, CK5 and CK14 peak values from line profiles are shown in box plots with inner and outlier points and mean. n(ctrl)=30; n(CK5 KD)=57; n(CK14 KD)=33. \*\*\*P<0.001 (paired ttest). (ttest, two tailed, equal variance).

(C) Western blot analyses on cell lysates from ctrl and CK5- as well as CK14-depleted cell lines showed downregulation of SLUG and slight upregulation of E-cadherin upon loss of CK5, as detected by specific antibody. GAPDH was used as a loading control.

(D) Quantification of SLUG and E-cadherin Western blot experiments. Mean ( $\pm$ SEM) is shown; n = 3; \*P<0.05; n.s.= not significant (paired ttest).

(E) A hypothetical model for the role of CK5 in the differentiation of mammary epithelial cell lineages, possibly through the regulation of SLUG. Some of the markers involved in this study are shown as examples within specific cell populations. Note that in the interest of space, several markers are missing from the hypothetical model and that in this study we did not concentrate on the expression pattern of these markers in distinct differentiation phases of the mammary epithelial cell populations. See f.i. Böcker et al. (2002); Boecker and Buerger (2003); Villadsen et al. (2007); Boecker et al. (2018); Fu et al. (2020) for such studies. See also Figure S8.

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impacts the mechanical features of the basal layer but also leads to loss of protective basement membrane possibly through impaired maturation of myoepithelial cells.

#### Loss of CK5 leads to downregulation of SLUG

To further assess the mechanisms through which CK5 and CK14 could impact the differentiation of mammary epithelial cells, we analyzed the levels of SLUG, a master regulator of the mammary epithelial cell lineage determination and normal tubulogenesis (Nassour et al., 2012; Phillips et al., 2014). In CK5, and to a lesser extent in CK14-deficient cells, SLUG was downregulated (Figures 6C and 6D). In line with these observations, the level of E-cadherin, a known target for SLUG-mediated repression (Bolós et al., 2016), was slightly upregulated upon depletion of CK5 (Figures 6C and 6D). Furthermore, depletion of CK5 by specific siRNAs from MCF10A mammary epithelial cells led to similar results and, additionally, loss of  $\alpha$ -SMA by siRNA had an almost equal impact (Figures S8D–S8F), again indicating reciprocal regulation within the basal cell populations. It should be noted that long-term downregulation of CK5 in cell culture conditions leads to upregulation of some other cytokeratins, including CK6, indicating that loss of CK5 is compensated through an alternative mechanism. This is supported by the re-induction of the studied myoepithelial markers and upregulation of several cytokeratins in the long-passaged cell clones (Figure S9).

These data indicate that the loss of CK5 may lead to differentiation bias in the mammary progenitors through regulation of SLUG levels. How specifically CK5 impacts SLUG levels, needs to be assessed in future studies. A hypothetical model for CK5 in the regulation of mammary epithelial lineage differentiation and formation of an intact, functional ME layer is presented in Figure 6E.

#### DISCUSSION

Mammary myoepithelial cells are important for normal mammogenesis and organostructural homeostasis, and have additionally been shown to have tumor suppressive properties (Gudjonsson et al., 2002; Jones et al., 2003; Polyak and Hu, 2005; Sánchez-Céspedes et al., 2016). Absence of ME cells and BM penetration determines stromal invasion, and a gradual loss of ME markers has been suggested to concur with malignant transformation of intraductal epithelial cells with subsequent breakdown of the protective ME barrier (Hilson et al., 2009; Kalof et al., 2004; Rohilla et al., 2015; Russell et al., 2015; Werling et al., 2003; Zhang et al., 2003). However, the molecular mechanisms behind the maintenance of this suggested myoepithelial barrier function are still poorly understood.

In this study, our goal was to understand in more detail the mechanisms leading to compromised ME barrier function. For this, we used a comparative canine model with immunohistochemical serial stainings for CK5, CK14, and  $\alpha$ -SMA. With these markers we were able to demonstrate some alterations in the basal/ME layer, in non-invasive intraductal proliferations of increasing grade of malignancy (UDH, ADH/L-G DCIS, I-G DCIS) by quantitatively determining their DAB chromophore staining intensity values in the TDLU and efferent inter-lobular ducts (Figures 1C and S2B). Of these markers, only  $\alpha$ -SMA displayed spatially statistically different expression patterns within the normal mammary ductal segments (Figure S1B). This result is in line with previous reports which have made semi-quantitative estimates on differences between the lobular alveolar/ductal and extralobular ductal compartments in the expression of some basal/ME markers in normal human mammary epithelium (e.g. Chen et al., 2015; Foschini et al., 2000; Pusztaszeri, 2010). This observation is possibly connected to the compartmentalization of the mammary epithelial structures into the intralobular functional alveolar and proliferative ductal zones and the extralobular efferent ductal system (Böcker et al., 2002; Rønnov-Jessen et al., 1996; Pusztaszeri, 2010). The impact of the differential composition of the surrounding intra- and extralobular stromal tissue on the segment-specific expression pattern should be further investigated.

Furthermore, we showed that the expression patterns of CK5 and  $\alpha$ -SMA of the basal/ME layer in the canine intralobular terminal ductal segments and the extralobular terminal/interlobular ductal segments undergo modest changes upon non-invasive intraductal proliferations (Figures 1C and S2B). In the extralobular terminal/interlobular ductal segment, the expression of these markers appears to slightly decrease already in the intermediate-grade DCIS prior to the invasive stage and is eventually lost at invasion (Figures 1A–1C). What is the biological significance or whether the cytoskeletal markers respond to non-invasive intraductal epithelial proliferations needs further studies.

 $\alpha$ -SMA is lost from the basal/ME layer prior to invasion (Russell et al., 2015). A recent study suggested that expression of  $\alpha$ -SMA, mediating the contractile properties of the ME layer, is essential for the mechanical barrier





function of ME cells against an invasion of transformed epithelial cells (Sirka et al., 2018). Our experiments showed that depletion of  $\alpha$ -SMA from the basal layer in 3D cultures led to abnormal mammosphere morphology (Figure 2), supporting the observation that contractility and mechanical features of the ME layer are crucial for the maintenance of normal mammary organostructure. Interestingly, loss of CK5 and CK14 from the basal layer resulted in similar, abnormally large and irregular 3D morphology (Figures 2, S5B, and S5C). Since cytokeratins are important for the mechanical features of epithelial cells, as also shown in our cell indentation experiments (Figures 3A–3D), KD of CK5 and CK14 could lead to abnormal compliancy of the basal layer and in this way advance such drastic morphological defects in the 3D mammospheres.

CK5 and CK14 are expressed in mammary stem and progenitor cell populations (Böcker et al., 2002; Boecker et al., 2018; Lee et al., 2012; Villadsen et al., 2007). However, their functions in the progenitors are not properly understood. Here, we show that KD of CK5, and to a lesser extent of CK14, affected the lineage commitment of the mammary progenitors: CK5-depleted CD49f<sup>+</sup> EpCAM<sup>-</sup>/basal progenitor-enriched cells showed impaired maturation into contractile ME cells, which was indicated by lower levels of  $\alpha$ -SMA, vimentin, SMMHC and calponin 1, and a concurrent increase in the expression of CK18 (Figures 3 and S6). Loss of CK14 had in our studies only a slight effect on these ME cell markers (Figures 3 and S6) and, as it is known to heterodimerize with CK5, this slight effect could possibly also go through CK5.

Additionally, ME-specific cell-cell junction proteins P-cadherin and Dsg3 were downregulated upon loss of CK5, causing deficiency in the maintenance of intact epithelial structures (Figure 4). Spatially selective expression of P-cadherin in mammary ME layer is required for the integrity of epithelial tissues and normal mammary architecture, and it has been shown that KD of P-cadherin from the ME cells compromises the barrier function of this cell layer (Idoux-Gillet et al., 2018; Sirka et al., 2018; Vieira et al., 2014). Furthermore, Dsg3 has been shown to co-localize with CK5 and CK14 and is linked to mechanotransduction through Ecadherin complex, indicating a role for this cell junction protein both in the maintenance of epithelial integrity and in adjustment of mechanical resistance in response to increasing external forces (Uttagomol et al., 2019; Vielmuth et al., 2018). Loss of CK5 from the basal progenitors thus affects the compliance, contractility, and integrity of the epithelial junctions, clearly leading to loss of ME barrier function. Interestingly, loss of  $\alpha$ -SMA from the ME cells also led to downregulation of CK5, vimentin, and Dsg3, indicating a regulatory feedback loop mechanism in between CK5 positive stem/progenitor cells and mature ME cells (Figures 4G, S6F and S6G). Furthermore, as CK5 KD led to impaired production of basement membrane protein laminin-5 around the 3D mammospheres (Figures 6A,6B, and S8A–S8C), the results indicate that CK5 not only impacts the mechanical features of the basal layer but also affects barrier function through the regulation of basement membrane, which is deposited by mature myoepithelial cells.

As  $\alpha$ -SMA has been suggested to be the main protein to mediate the contractile potential of ME cells (Haaksma et al., 2011), we expected that loss of the mature ME cell phenotype would lead to a cell type which exerts less forces on its environment. However, actomyosin-mediated cell-substrate forces were slightly increased, as detected by traction force microscopy with single cells and also with monolayers (Figures 5,S7C, and S7D). This may be explained by the lower levels of these specific cytokeratins as well as the subsequent lower levels of vimentin, since intermediate filaments have been indicated to play a role in the co-regulation of actomyosin forces through their association with cell adhesion sites (Bordeleau et al., 2010, 2012; Jiu et al., 2017). The KD phenotype could thus exert uncontrolled forces on the underlying substrate. Alternatively, weakened cell-cell junctions and the appearance of more prominent cell-substrate adhesions in the KD cell lines could result in redistribution of cellular forces more toward the underlying substrate. Whether this has an impact on cellular motility needs to be further assessed in the future.

Finally, the loss of mature ME cell phenotype upon CK5 KD was associated with slightly higher expression of luminal marker CK18 (Figures 3 and S6), indicating a differentiation bias towardthe luminal cell type. CK5-deficient cells also expressed significantly lower levels of the transcriptional repressor SLUG (Figures 6C, 6D, and S8D–S8F). SLUG has been shown to determine the lineage specific differentiation of mammary epithelial cells and is co-localized in a subpopulation of basal cells together with CK5, P-cadherin, and CD49f (Nassour et al., 2012). In line with our observations, SLUG-deficient cells have been shown to over-express higher levels of markers linked to luminal lineage, such as CK8, CK18, and ER (Nassour et al., 2012). SLUG-deficient adult mice display abnormal mammary epithelial cell lineage differentiation with increased expression of luminal markers in the basal layer and hyperplasia of luminal cells (Phillips et al., 2014). Supporting that, our studies showed that CK5 KD, and to a lesser extent CK14 KD cell lines, showed increased





expression of CK18 (Figure 3). In breast cancer, increased CK18 expression has been linked with inhibition of apoptosis, increase in the expression of CK8 and adhesion proteins as well as decrease in vimentin levels (Aiad et al., 2014; Bozza et al., 2018; Bühler and Schaller, 2005; Iyer et al., 2013; Schaller et al., 1996; Weng et al., 2012). Moreover, we observed that E-cadherin, a target for SLUG-mediated repression (Bolós et al., 2016), was upregulated upon depletion of CK5 (Figures 6C,6D,S8D, and S8E). CK5 KD cells were also growing slower and a similar phenotype has been observed in SLUG-deficient cells (Nassour et al., 2012). As SLUG clearly plays a role in the maintenance of basal-like state and represses luminal lineage differentiation, loss of CK5 could conceivably cause the differentiation bias via regulation of SLUG. The exact mechanisms through which CK5 impacts SLUG levels needs to be further studied in the future.

In conclusion, our findings support the previous studies that have underlined the importance of basal myoepithelial cell layer as a barrier that is eventually lost prior to the invasive stage. Our data showed that CK5 loss plays a major role in the disruption of this myoepithelial layer leading to defects in basement membrane formation. Downregulation of CK5 and consequent loss of SLUG led to epithelial cell differentiation bias with subsequent defects in the maturation of myoepithelial cells and a shift toward the CK18-positive luminal epithelial cell type. The reciprocal interactions of these proteins should also be assessed in more detail in the future.

#### LIMITATIONS OF THE STUDY

Although this study shows an interesting link between cytokeratin 5 and SLUG expression, possibly playing a role in the differentiation of specific mammary epithelial cell populations, this work does not provide any information on the molecular mechanisms behind this interconnection. The role of CK5 in the regulation of SLUG levels clearly needs further studies in the future. Also, the technical challenges in the 3D mammo-sphere antibody-stainings limited these studies. Furthermore, the amount of canine patient samples, related to Figure 1, was very limited.

#### **Resource** availability

#### Lead contact

Further information and requests for resources and reagents should be directed to and will be fulfilled by the Lead Contact, Sari Tojkander (sari.tojkander@helsinki.fi).

#### Materials availability

This study did not generate new unique reagents. Materials are available on request.

#### Data and code availability

This study did not generate any unique datasets or code. All raw data is available on request.

#### **METHODS**

All methods can be found in the accompanying Transparent methods supplemental file.

#### SUPPLEMENTAL INFORMATION

Supplemental information can be found online at https://doi.org/10.1016/j.isci.2021.102413.

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#### **AUTHOR CONTRIBUTIONS**

V.D. designed experiments, has collected and diagnosed canine mammary samples, performed IHC staining on FFPE canine mammary samples, collected and analyzed the data for determination of DABi, isolated canine mammary cells with FACS, analyzed and interpreted the data, and prepared Figures 1 andS1–S3 and the manuscript; K.R. has performed siRNA experiments, 3D Matrigel experiments, IF stainings and WBs, contributing to Figures 2, 3, 4, 5, 6, and S3G–S9 and participated in writing the manuscript; S.C. has performed KD cell lines, isolated human mammary cells with FACS, and performed initial WB analyses for Figures 3 and S4; A.A. has participated in the 3D mammary epithelial culture experiments, and Western blot experiments in Figures 2F, 4F, 6A, S5B, S5C, S7A, and S8A–S8C. N.S and R.K. have co-analyzed the traction force data. J.J.V.-D. and M.Ö. have performed AFM experiments and analyses of the data. P.B. has been involved in canine mammary tumor sample collection and provision of clinical patient data; A.S. has been involved in canine mammary tumor sample collection and has provided pathological expertise; S.T. has designed experiments, performed TFM experiments and co-analyzed the data, prepared figures and co-written the manuscript.

#### **DECLARATION OF INTERESTS**

The authors declare no competing interests.

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# iScience Article

iScience, Volume 24

# Supplemental information

# Cytokeratin 5 determines maturation

# of the mammary myoepithelium

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### 32 TRANSPARENT METHODS

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# 34 Histological samples of canine mammary tissues

Formalin-fixed paraffin-embedded tissue samples of female canine mastectomy cases (n = 7) with non-infiltrative intraductal proliferative lesions were retrieved from the pathologic-diagnostic archive (Section of Veterinary Pathology, University of Helsinki). Normal mammary gland from the same individual was prerequisite. Respective epidemiological data such as breed, sexual status and age at diagnosis were collected (Fig. S2A).

Histomorphology was reviewed from 4 µm thick hematoxylin and eosin stained tissue sections. 40 Classification criteria of woman for usual ductal hyperplasia (UDH), atypical ductal hyperplasia 41 (ADH) and ductal carcinoma in situ (DCIS) were adopted for canine female non-infiltrative 42 43 intraductal proliferative lesions (The Consensus Conference Committee, 1997; Collins et al., 2012; Ferreira et al., 2012; Goldschmidt et al., 2011; Mouser et al., 2010; Schnitt et al., 2012b; Simpson et 44 45 al., 2012). Canine lesions classified as ductal displayed E-cadherin positivity, and invasion was determined as absence of myoepithelial cells and/or extension through the basement membrane 46 47 (Chocteau et al., 2019; Goldschmidt et al., 2011; Ressel et al., 2011; The Consensus Conference Committee, 1997). Canine UDH applied to intraductal proliferation of small epithelial cells with 48 49 hyperchromatic nuclei, scant cytoplasm and little nuclear or cellular pleomorphism. Subcategorization as mild, moderate or florid based on the amount of hyperplastic cells forming 50 51 epithelial bridges and irregular fenestrations filling the ductal lumen. The intraluminal cells displayed positivity for low and/or high molecular weight cytokeratins. Lesions with increased cellular atypia 52 corresponding to low-grade DCIS, but with only partial involvement of the ductal lumen and limited 53 extension classified as ADH. DCIS applied to intraductal epithelial proliferations of increasing 54 cellular atypia allowing categorization into low, intermediate and high nuclear grade DCIS. 55

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# 57 Tissue section immunohistochemistry

Formalin-fixed paraffin-embedded female canine mammary tissue samples were prepared into 4 µm 58 thick serial sections on Menzel Superfrost Plus Adhesion microscope slides (Cat. J1800AMNZ, 59 Thermo Fisher Scientific) and baked at 37 °C o/n. The slides were deparaffinized and rehydrated 60 automated (Leica AutoStainer XL, Leica Biosystems) in xylen and alcohol series before antigen 61 retrieval in a PT Module (LabVision UK Ltd) for 20 min at 99 °C in prewarmed 0.01 M citrate buffer 62 (pH 6.0). After cooling down to RT, the slides were blocked 10 min with 3% hydrogen peroxide in 63 PBS, and rinsed twice with TBS + Tween (Cat. P1379, Sigma). For staining, the BrightVision Poly-64 HRP-Anti Ms/Rb/Rt IgG Kit (Cat. DPVO110HRP, Immunologic) was used at RT with minor 65

modifications to manufacturers protocol. Primary mouse antibodies were incubated 1 h at RT and 66 include anti-CK5 (Clone XM26, Cat. 17130, Abcam, dilution 1:75), anti-CK14 (Clone LL002, Cat. 67 7800, Abcam, dilution 1:300), anti-CK18 (Clone Ks18.04, Cat. 61028, Progen, dilution 1:300), anti-68 α-SMA (Clone 1A4, Cat. M0851, Dako, dilution 1:600) and primary rabbit anti-E-cadherin (Clone 69 24E10, Cat. 3195, Cell Signaling Technology, dilution 1:500). For visualization, 3.3'-70 diaminobenzidine-tetrahydrochloride (Cat. VWRKBS04-110, Immunologic) was applied for 5 min. 71 72 Harris Hematoxylin (Cat. HX57998853, Merck) was used for 10 sec to counterstain. The slides were dehydrated and mounted with Pertex (Cat. 00811, Histolab). To ensure equal preparation and staining 73 74 conditions for all the tissue slides, the protocol was implemented on as one batch in one same run. Adjacent normal tissue was internal positive control. From negative controls the antibody was omitted 75 and antibody diluent only was used. Stainings were performed twice. 76

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# 79 Determination of cellular DAB chromogen precipitate intensity

80 For determination of color-intensity of basal myoepithelial cytoplasmic 3.3'-diaminobenzidinetetrahydrochloride chromogen precipitate (i.e. DAB staining intensity, DABi), digital images were 81 82 generated from IHC stained (α-SMA, CK5 and CK14) female canine mammary tissue slides using a Pannoramic 250 FLASH II digital whole-slide scanner (3DHISTECH Ltd., Budapest, Hungary) with 83 a VCC-FC60FR19CL camera (CIS Corporation, Japan) and LS-6 pulsed Xenon light source 84 (Excelitas Technologies Corp., Waltham, MA, USA). Extended focus was applied with the following 85 86 settings: Focus distance field of views: 8; Focus levels: 7; Step size (0.2 µm): 5. Resolution of the scanner was 0.24 µm/pixel with a 20x/NA 0.8 objective. To ensure subjection to same imaging 87 conditions, tissue slides were scanned as one batch. For analysis, representative areas of the features 88 (normal, UDH, ADH, DCIS) from each case were imaged at 69.12x magnification using the 89 Pannoramic Viewer Software version 1.15.4 (3DHISTECH Ltd., Budapest, Hungary). 90

91 Using Fiji ImageJ Software version 1.51 (U. S. National Institutes of Health, Bethesda, Maryland, USA) the blue wavelength band was separated from TIFF (Tagged Image File Format) image and 92 inverted to 8-bit resolution grey-scale image with intensity values ranging from 0 = black to 255 =93 white. DABi was determined from individual basal/myoepithelial cells by freehand drawing the 94 boundary of each individual cellular cytoplasmic area (i.e. region of interest, ROI) to be quantified. 95 Inclusion criteria for cells to be analyzed included basal location between the basement membrane 96 and the luminal epithelial cell layer as well as the presence of a nucleus. Manual work allowed 97 precision excluding from the analysis such features as cellular nuclei, overlapping cells and cellular 98 areas showing only cytoplasm without nuclei as well as other positive reactions for the given antibody 99

(e.g. for α-SMA adjacent myofibroblasts and blood vessels) which would otherwise interfere with the
results. DABi was collected as integrated density value, being the product of ROI area and mean pixel
value. Mean pixel value is determined as the sum of the grey values of all the pixels in the ROI
divided by the number of pixels. DABi values from the intralobular terminal ductal and the
extralobular terminal ductal/interlobular ductal segments were collected and analyzed as distinct
entities according to their anatomical and physiological differences (Rønnov-Jessen et al., 1996). The
used protocol is summarized in Fig. S3B.

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## 109 Isolation of canine mammary primary organoids

Canine normal mammary tissue samples (c. 5 x 10 mm) were collected aseptically into DMEM/F12 110 medium (Cat. 31330-038, Gibco) from removed tissue (sterilized female, 9 years-of-age, mixed breed 111 of Belgian Sheepdog and German Shepherd Dog) at canine mammary gland surgery (Veterinary 112 Teaching Hospital, University of Helsinki), and kept at 4 °C until macrodissection to remove adherent 113 fat and surplus stroma (LaBarge et al., 2013). The medium was supplemented with 0.25 µg/ml 114 Amphotericin B (Cat. A2942, Sigma), 100 U/ml Penicillin and 100 µg/ml Streptomycin (Cat. 15140-115 116 122, Gibco) as well as 10% FBS (Cat. 10500-064, Gibco). Mammary tissue next to the area of sample origin was collected simultaneously into 10% neutral-buffered formalin to confirm histomorphology. 117 118 After macrodissection, the remaining tissue was cut into 3-4 mm slices for digestion in pre-warmed (37 °C) DMEM/F12 medium supplemented with antibiotica and FBS (as above), 10 µg/ml Insulin 119 120 (Cat. I6634, Sigma), 200 U/ml Collagenase (Cat. C0130, Sigma), and 100 U/ml Hyaluronidase (Cat. H3506, Sigma) with o/n incubation in a 37 °C water bath using gentle agitation. Digestion was 121 monitored by microscopy using a camera (Canon EOS 600D) attached to an inverted microscope 122 123 (Olympus CKX41).

Digested material was centrifuged at 600 x g 5 min to separate remaining fat etc. (LaBarge et al., 2013). A 40 µm cell strainer was used to collect the size-differentiated organoid fractions. Media supplemented with antibiotics and FBS (as stated above) was applied for rinsing. The collected material was pelleted at 80 x g 30 s and aliquoted into freezing medium of 50% DMEM/F12 plus 44% FBS plus 6% DMSO (Cat. D2650, Sigma) (StemCell Technologies, 2012) and stored at -130 °C until fluorescence-activated cell sorting (FACS).

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# 131 Fluorescence activated cell sorting of canine mammary epithelial cells

For FACS, the canine organoid fractions were incubated at 37 °C in a humidified 5%  $CO_2$  cultivator in DMEM/F12 supplemented with antibiotica and FBS (as stated above), 15.0 µg/ml Insulin (Cat.

I6634, Sigma), 10.0 ng/ml EGF (Cat. E9644, Sigma) and 0.6 µg/ml Hydrocortisone (Cat. H0888, 134 Sigma) for 2-3 d. Differential trypsinization to remove fibroblasts was applied prior generating single-135 cell suspension for FACS (LaBarge et al., 2013). Single cell suspension of 1x10<sup>6</sup> cells/ml was 136 generated into HBSS + 1% BSA from canine primary culture using incubation with TrypLE Express 137 (Cat. 12604-013, Gibco) and DNase I (Cat. 11284932001, Roche). Cells were incubated with anti-138 CD24-Alexa Fluor 405 (Clone ML5, Cat. NB100-77903AF405, Novus Biotechne, 0.7mg/ml) at 4 °C 139 for 45 min in the dark and washed with HBSS + 1% BSA three times. After the last wash, the cells 140 were re-suspended into 500  $\mu$ l FACS-buffer consisting of PBS + 0.5% BSA followed with a 10 min 141 incubation with a live/dead discriminating stain (Cat. L34975, Invitrogen/Thermo Fisher Scientific). 142 A FACSAria II (BD Biosciences) instrument was used to sort the cells with red (633 nm) and violet 143 (405 nm) lasers at sheat pressure 45.00 and with a 85 µm nozzle. The data was collected using the 144 FACSDiva Software version 8.0.2 (BD Biosciences). Dead cells, non-single cells and CD24<sup>neg</sup> cells 145 146 were excluded. The cells were isolated into growth medium (as stated above) in tubes coated by incubation with HBSS + 3% BSA for 2 h under UV-radiation at RT. To determine the sorting purity, 147 148 the isolated cell population was stained for immunofluorescence imaging prior downstream assay (Fig. S2F). FACS data was analyzed and the graphs generated with the FlowJo Software version 149 150 10.4.1 (BD Biosciences).

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# 152 Cell culture and siRNA transfections

Human breast epithelial cell line MCF10A cells (ATCC) were maintained in DMEM/F12 media
supplemented with 5% Horse Serum (Cat. 26050088, Gibco), 20 ng/ml EGF (Cat. E9644, Sigma),
0.5 mg/ml Hydrocortisone (Cat. H0888, Sigma), 100 ng/ml Cholera toxin (Cat. C8052, Sigma), 10
µg/ml Insulin (Cat. I6634, Sigma) and 100 U/ml Penicillin and 100 µg/ml Streptomycin (Cat. 15140122, Gibco). Cells were cultured in an incubator with 5% CO<sub>2</sub> at 37 °C.

For siRNA transfections in 2D, MCF10A cells were plated on 35 mm dishes at a density of 5000 cells 158 per well. The following day, cells were transfected with 50 nmol siRNA (human CK5: ON-159 TARGETplus KRT5 SMART pool siRNA [L-011067-00-0005]; human CK14: ON-TARGETplus 160 KRT14 SMART pool siRNA [L-010602-00-0005]; and human SMA: ON-TARGETplus ACTA2 161 162 SMART pool siRNA [L-003450-00-0005]; negative control: ON-TARGETplus Non-targeting siRNA [D-001810-01-05] from Dharmacon) using Ribojuice siRNA transfection reagent (Cat. 163 71115-3, Millipore) according to the manufacturer's instructions. On day 4., cells were processed for 164 Western blotting or microscopy. Canine primary cell transfections in 2D conditions were performed 165 as with human MCF10A cells. For canine cells, pool of custom-made siRNAs were used: canine 166 CK5: 5' UAACUCUUGAAACUCUUCCUU 3', 5' AACUCUUGAAACUCUUCCGUU 3', 5' 167

ACUCUUGAAACUCUUCCGGUU 3'; canine CK14: 5' UAGUCCUUGGUCUCAGCGGUU 3',
5' UCUCAGAGCGUUCAUUUCCUU 3', UCCACGUUGACAUCUCCGCUU 3'; canine ACTA2
5' UACUUCAAGGUCAGGAUCCUU 3', 5' UCUAUCGGGUACUUCAAGGUU 3', 5'
AUGAUGCCGUGUUCUAUCGUU 3' (Dharmacon). In 3D Matrigel cultures of human and canine
mammary epithelial cells, siRNAs (50 nmol) were applied after formation of mammospheres (day 5)
and incubated for 2 weeks. siRNAs were added to cells every 4 days.

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## 175 Fluorescence cell sorting of MCF10A cells

176 The FACS protocol from Abcam was followed in this experiment. Briefly, the cells were harvested by trypsinization. Disassociated cells were resuspended at a concentration of approximately  $1 \times 10^{6}$ 177 cells/ml in ice cold PBS, 10% FCS, 1% sodium azide. The cells were fc blocked using Human Fc 178 block (Cat. 564219, BD Biosciences) following manufacturer's instructions. To remove unbound fc 179 180 block, the cells were washed 3 times by centrifugation at 400 g for 5 min and resuspended in ice cold PBS. 5 µg/ml of the primary antibodies anti-CD49f-FITC (Clone GoH3, Cat. 561893, BD 181 182 Biosciences) and anti-EpCAM-APC/Cy7 (Clone G8.8, Cat. 118217, Biolegend) were added to the cells and incubated for 1 h at 4°C in the dark. The cells were washed as described in the previous 183 184 steps and resuspended in ice cold PBS, 10% FCS, 1% sodium azide. The cell suspension was immediately stored at 4°C in the dark. The cells were sorted using a FACSAria II (BD Biosciences) 185 instrument at sheat pressure 45.00 with a 85 µm nozzle and the blue (488 nm) as well as red (633 nm) 186 lasers. The flow cytometer settings adjustments and subsequent data collection was done using 187 FACSDiva Software version 6.1.3 (BD Biosciences). Compensation was performed using unstained 188 cells and single stained control cells. The data generated was analyzed using FlowJo Software version 189 10.6.1. 190

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# 192 Lentiviral-based RNA interference

Lentiviral RNA interference was done as described in Cattavarayane et al., 2015. The packaging 193 vector containing the desired shRNA construct was received from SIGMA. The helper plasmids 194 pMD2.G (Plasmid 12259), and pMDL g/p RRE (Plasmid 12251) and pRSV-Rev (Plasmid 12253) 195 were provided by Addgene. DNA transfections were performed by using Lipofectamine 2000 196 (Invitrogen). The packaging vector: pVSVG : pMDL g/p RRE : pRSV-Rev were used in 3:1:1:1 197 proportion and the total DNA used for transfection was 20  $\mu$ g (15 $\mu$ g :: 2.5  $\mu$ g : 830 ng : 830 ng : 830 198 ng). Nearly confluent (70-80%) 293 T cells were grown on Corning CellBind (Cat. 3296) 6 well plate 199 for transfections. 24 hrs post transfection, media containing the transfection mix was removed 200 carefully and 1 ml of fresh media was added. The media supernatant containing the viral particles 201

was collected every 12 hrs for 3–4 days and stored at 4 °C. The viral supernatants were pooled and centrifuged at 1000 rpm for 5 minutes and filtered through a 0.44  $\mu$ m filter. The cells were infected using viral supernatant for 24 hrs in the presence of 4  $\mu$ g/ml polybrene (Cat. 107689, Sigma) and the infected cells were selected using puromycin (4  $\mu$ g/ml, Sigma). The puromycin selected cells were expanded and stored by freezing in 90% FCS, 10% DMSO (Sigma) at -130 °C.

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# 208 Western blotting

Cells were washed with PBS and lysed in 1% Triton X-100/PBS, containing protease and phosphatase 209 inhibitors (Cat. 539131 and Cat. 539131, Calbiochem). Protein concentrations were measured with 210 Qubit® Protein Assay Kit (ThermoFisher Scientific). Sample loading buffer, 4x LSB-DTT, was 211 added to lysates and samples were boiled for 5' before loading in SDS-PAGE gels (Bio-Rad). Wet 212 transfer (Bio-Rad) with Immobilon-P Membrane, PVDF filter (Millipore) was used for blotting. 213 Mixture of 5% BSA and 5% milk was used for blocking for at least 1 h. Following antibodies were 214 215 used for detection of specific proteins: mouse anti-CK5 (Clone XM26, Cat. ab17130, Abcam), mouse anti-CK6 (Clone B-7, Cat. sc-514520, Santa Cruz), mouse anti-CK14 (Clone LL002, Cat. ab7800, 216 217 Abcam), rabbit anti-CK18 (Polyclonal, Cat. ab24561, Abcam), mouse anti-CK19 (Clone BA-17, Cat. ab7755, Abcam), mouse anti-Pan-Keratin (Clone C11, Cat. 4545, Cell Signaling Technology; 218 Recognizing CK4, 5, 6, 8, 10, 13 and 18), mouse anti-SMA (Clone 1A4, Cat. A5228, Sigma), rabbit 219 anti-E-cadherin (Clone 24E10, Cat. 3195, Cell Signaling Technology), rabbit anti-P-Thr18/Ser19-220 MLCII (Clone Thr18/Ser19, Cat. 3674, Cell Signaling Technology), mouse anti-calponin1 (Clone 221 CALP, Cat. sc-58707, Santa Cruz), rabbit anti-Vimentin (Clone D21H3, Cat. 5741, Cell Signaling 222 Technology), mouse anti-Dsg3 (Clone 5H10, Cat. sc-23912, Santa Cruz), rabbit anti-P-cadherin 223 (Clone C13F9, Cat. 2189, Cell Signaling Technology), rabbit anti-Slug (Clone C19G7, Cat. 9585, 224 Cell Signaling Technology) and mouse anti-GAPDH (Clone GAPDH-71.1, Cat. G8795, Sigma). 225 Anti-mouse or -rabbit HRP-linked secondary antibodies (Cell Signaling Technology) in 1:3000 226 dilution and Western HRP substrate (Cat. WBLUR0100, Luminata<sup>TM</sup>Crescendo, Millipore) were 227 used for chemiluminescence detection of the protein bands. Images were taken with Fujifilm LAS-228 3000 Imager using autoexposure-setting that choose the optimal exposure time for each blot. 229

230

## 231 Immunofluorescence microscopy

Cells were cultured on coverslips and washed with PBS prior to fixation with 4% PFA.
Permeabilization was performed with 0.1% Triton X-100 in TBS for 5' and cells on coverslips were
moved to 0.2% Dulbecco/BSA. The following primary antibodies were used for stainings in 1:50
dilutions: mouse anti-CK5 (Clone XM26, Cat. ab17130, Abcam), mouse anti-CK14 (Clone LL002,

Cat. ab7800, Abcam), rabbit anti-P-Ser18/Thr19-MLCII (Clone Thr18/Ser19, Cat. 3674S, Cell 236 Signaling Technologies), rabbit anti-E-cadherin (Clone 24E10, Cat. 3195, Cell Signaling 237 Technologies), mouse anti-vinculin (Clone hVin-1, Cat. V9264, Sigma), and mouse anti-Dsg3 (Clone 238 5H10, Cat. sc-23912, Santa Cruz), mouse anti-Laminin-5 (Clone D4B5, Cat. MAB19562, Sigma-239 240 Aldrich). Alexa Fluor  $\alpha$ -rabbit 488 and  $\alpha$ -mouse 568 (Life TechnologiesTM) secondary antibodies were used to detect the primary antibodies. Actin cytoskeleton was detected with Alexa-488-, -568-241 and -647-Phalloidins in 1:200 dilution (Life TechnologiesTM) and DNA with DAPI (Life 242 TechnologiesTM). DABCO/Mowiol was used for mounting. Images were acquired with Leica 243 DM6000 and Leica DM5000 upright fluorescence wide field microscopes equipped with a 244 Hamamatsu Orca-Flash4.0 V2 sCMOS camera. 245

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# 247 **3D mammosphere cultures**

248 Matrigel, ECM gel from Engelbreth-Holm-Swarm murine sarcoma (Sigma E1270) was prepared according to manufacturer's instructions. MCF10A cells were trypsinized and approximately 5000 249 250 cells in F12 media were seeded on top of the Matrigel-coated eight-chamber slides. Formation and morphology of spheres were monitored frequently under a light microscope. After two weeks, 3D 251 252 structures were fixed with 2% PFA and diameter of the structures were calculated from bright field images using ImageJ. For IF, cells were permeabilized with 0.25% Triton-X in PBS for 10 min. 253 Unspecific binding sites were blocked with IF buffer (0.1% BSA, 0.2% Triton-X, 0.05% Tween in 254 PBS) supplemented with 10% goat serum at RT for 1 h. Primary antibodies in blocking solution were 255 incubated o/n at 4 °C and secondary antibodies at RT for 30 mins. Cytoskeletal structures were 256 stained with Phalloidin and nucleus with DAPI. Samples were mounted with Mowiol/DABCO. The 257 258 IF stainings of 3D spheroids were performed as in https://brugge.med.harvard.edu/protocols with minor modifications. Imaging stacks was performed with Leica TCS SP5. Objective used: HCX PL 259 APO 20x/0,7 Imm Corr (water, glycerol, oil) Lbd.bl. 260

261

# 262 Determination of cell elastic modulus by atomic force microscopy

The elastic moduli of MCF10A, CK5 KD, and CK14 KD cells were determined by cell indentation experiments using a MultiMode 8 atomic force microscope (AFM) with a NanoScope V controller and a closed-loop PicoForce scanner (Bruker, Santa Barbara, CA). The cells, grown on coverslips coated with LN111, were mounted in the AFM and indented with triangular MLCT probes (Bruker) with nominal tip radius of 20 nm and spring constants about 0.05 N/m, obtained by the thermal noise method (Hutter and Bechhoefer, 1993). The deflection sensitivity was measured on a hard, freshly

cleaved mica surface just before the experiments with cells. Cell indentation was carried out with the 269 AFM operating in Force Volume mode (contact mode), collecting indentation force curves in 40  $\times$ 270 40  $\mu$ m<sup>2</sup> areas, the so-called force maps, which were divided into 16 × 16 pixels. A force curve was 271 recorded at each pixel of the force map (that is, 256 force curves per force map) by indenting the cells 272 at 4 µm/s rate, with maximum applied forces typically in the range of 1.0–1.5 nN. The experiments 273 were carried out in PBS at 37 °C (Thermal Applications Controller, Bruker). The elastic modulus of 274 the cells were obtained by fitting the approach indentation curves with the Sneddon (conical indenter) 275 model (Sneddon, 1965), using the software NanoScope Analysis 1.5 (Bruker). Between 440 and 750 276 indentation curves from 2 or 3 different  $40 \times 40 \ \mu\text{m2}$  force maps, corresponding to about 8 to 12 277 cells, were analyzed for each cell type. The indentation depth was typically in the range  $1.5 - 2.5 \,\mu\text{m}$ . 278 The corresponding elastic modulus histograms were fit with three Gauss distributions using OriginPro 279 2020 software. 280

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# 282 Traction force microscopy

283 For traction force microscopy experiments in the single cell, the cell doublet, or the monolayers, cells were plated on elastic collagen-1-coated polyacrylamide (PAA) gel substrates with a stiffness 284 285 (Young's Modulus/elastic modulus) of 4 kPa (single cell) or 6.3 kPa (cell doublet) and incubated for 2-4 h prior to imaging. Gel substrates were surface-coated with sulfate fluorescent microspheres 286 (Invitrogen, diameter 200 nm) before coating with collagen-1. Single cells together with the 287 underlying microspheres were imaged at multiple locations with 3I Marianas imaging setup 288 containing a heated sample chamber (37 °C) and controlled CO<sub>2</sub> (3I intelligent Imaging Innovations, 289 Germany). 63x/1.2 W C-Apochromat Corr WD=0.28 M27 objective was used. Following live cell 290 imaging, the cells were detached from the substrates with 10x Trypsin (Lonza Group) and a second 291 set of microsphere images was taken to serve as reference images. Displacement maps for 292 microspheres were achieved by comparing the reference microsphere images to the experimental 293 images and by knowing the cell-induced displacement field, substrate stiffness (4 kPa), and a manual 294 trace of the cell boundary, we computed the cell-exerted traction fields by using Fourier Transform 295 Traction Cytometry (Krishnan et al., 2009; Tolić-Nørrelykke et al., 2002). Root mean squared 296 magnitudes were computed from the traction fields. For the measurement of intercellular stresses, we 297 utilized cell doublets. These assays were performed at 6.3 kPa PAA dishes, with an optical resolution 298 of 63x and analyzed at a spatial resolution of  $\sim 1 \mu m$ . 299

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## **303 Colocalization analyses**

Colocalization of actin (Phalloidin) and Dsg3 at cell-cell junctions was analyzed in Fiji. Manually drawn areas of cell-cell junctions were used as regions of interest, ROI, and colocalization (in %) on this region was analyzed with FIJI colocalization analysis.

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# 308 Analyses of laminin-5 intensity from 3D spheroids

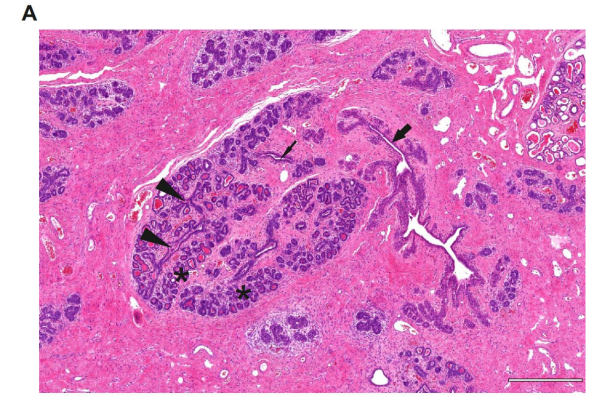
Raw data images from laminin-5 stainings of 3D mammospheres were imported to Fiji and line 309 profiles, starting immediately outside the spheroid structures, were drawn towards the cell center (Fig. 310 6A). 3-5 line profiles were drawn around one spheroid. Intensity maps of the stainings were also 311 performed to better demonstrate the differences in between samples. Numerical values from the line 312 profiles were exported to excel. Of these data, the third point from the line profiles mostly displayed 313 highest intensity values within the studied samples and was chosen for further analysis to compare 314 the laminin-5 intensities in between different samples/spheroids. t-test (two-tailed, equal variance) 315 was used to analyse the statistical significance of these peak values in between groups. 316

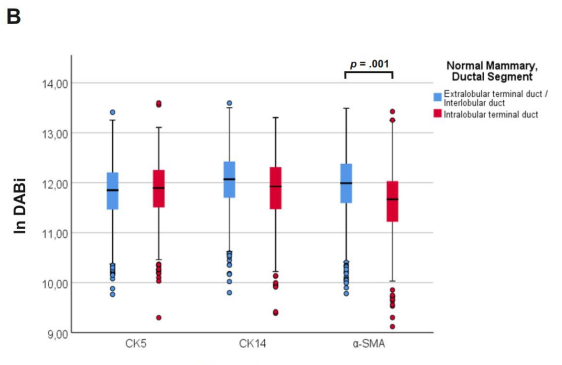
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### 318 Statistical analysis

Standard deviations and statistically significant difference between means of two groups (t-test) were 319 analyzed in Microsoft Excel 2013. Mann-Whitney-Wilcoxon rank-sum test was performed with 320 Origin 2018. For statistical analysis of DAB chromogen staining intensity (DABi), multilevel mixed 321 linear model was performed using the IBM SPSS Software version 25.0 (IBM Corp., USA). The 322 fixed main effects of age, gender, DABi biomarker group (i.e. CK5, CK14 and α-SMA), ductal 323 324 segment and pathologic-anatomic diagnosis (PAD) on the DABi-values were examined. The DABi biomarker group, PAD and the interaction between biomarker group and PAD were regarded as fixed 325 326 effects. Random intercepts for dog individual and ductal segment (i.e. intralobular terminal duct, extralobular terminal duct/interlobular duct) were used to account for the hierarchical structure of the 327 328 data. Thus, individual basal myoepithelial cells (i.e. observation unit) were nested (clustered) within 329 ductal segments and ductal segments nested within dogs. DABi-values were In-transformed for the 330 statistical analysis due to skewed distribution. Bonferroni's multiple comparisons correction was used when comparing more than 2 groups, that is DABi biomarker groups and PAD groups within and 331 332 between ductal segments. Pairwise comparison over all the individual companions was implemented. The level of significant was defined as p < 0.05. No statistical methods were used to predetermine 333 the DABi biomarker study sample size. 334





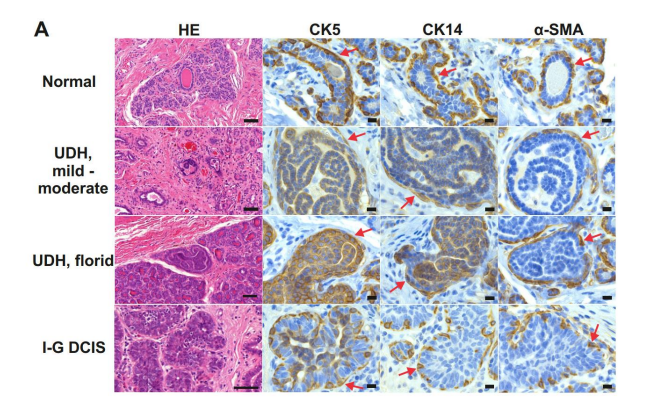


**Biomarker Group** 

# 339 Figure S1: Expression of basal myoepithelial biomarkers in normal mammary ductal segments,

Related to Figure 1. A) Histomorphology of the canine mammary terminal duct lobular unit (TDLU) 340 consisting of a lobule with acinar units (asterix) and intralobular terminal duct (wedge) as well as 341 extralobular terminal duct (thin arrow), leading into interlobular duct (thick arrow). Hematoxylin and 342 eosin (HE) staining. Bar 500 µm. B) Boxplot showing the difference in between the cellular In-343 transformed DAB intensity (In DABi) values for the basal myoepithelial biomarkers CK5, CK14 and 344 α-SMA in between the normal intralobular terminal ductal segment (CK5 n(cells)=612; CK14 n=816; 345 α-SMA n=875) and the extralobular terminal / interlobular ductal segment (CK5 n=1203; CK14 346 n=1469; α-SMA n=1765). Canine patient n=7. Black middle line within box represents median. 347 Height of box is interquartile range (IQR), representing 75<sup>th</sup> and 25<sup>th</sup> percentiles, respectively. 348 Whiskers represent the lowest and highest data within the 1.5 x IQR of the lower and upper quartiles, 349 respectively. Circles represent outliers. Linear mixed model with random intercepts for canine 350 individual and ductal segment was used with Bonferroni's multiple comparisons correction. The level 351 of significant was defined as p < 0.05. Only statistically significant mean differences are indicated. 352 Significant difference was determined between the normal ductal segments in the expression of  $\alpha$ -353 SMA (p = 0.001). 354

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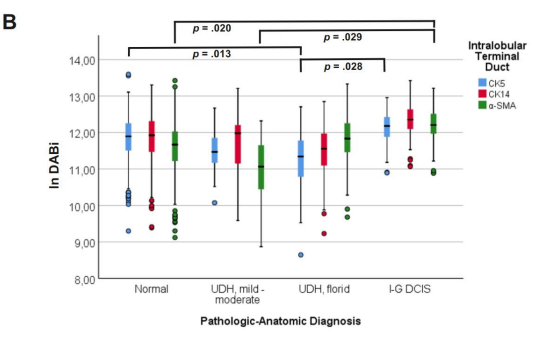


Figure S2. Expression of CK5, CK14 and  $\alpha$ -SMA in the intralobular terminal ductal segment upon intraductal hyperplasia, Related to Figure 1. A) Consecutive canine mammary FFPE tissue sections of normal, mild-to-moderate usual ductal hyperplasia (UDH, mild-to-moderate), florid usual ductal hyperplasia (UDH, florid) and intermediate-grade ductal carcinoma in situ (I-G DCIS) were stained using hematoxylin-eosin (HE, far left) and the basal myoepithelial markers CK5 (middle left), CK14 (middle right) and  $\alpha$ -SMA (far right). Representative images of the lesions are shown. Red

arrows indicate basal myoepithelial layer as distinct from intraluminal cellular hyperplasia. Bar for 364 HE in normal, mild-moderate UDH, florid UDH and I-G DCIS 50 µm, and in all IHC 10 µm. B) 365 Boxplot of the quantifications of intralobular terminal ductal cellular In-transformed DAB intensity 366 (ln DABi) values for normal CK5 n(cells)=612, CK14 n=816, α-SMA n=875; UDH mild-to-moderate 367 CK5 n=22, CK14 n=17, α-SMA n=29, UDH florid CK5 n=151, CK14 n=203, α-SMA n=258; I-G 368 DCIS CK5 n=105, CK14 n=184, α-SMA n=153. Canine patient n=7. Black middle line within box 369 represents median. Height of box is interquartile range (IQR), representing 75<sup>th</sup> and 25<sup>th</sup> percentiles, 370 respectively. Whiskers represent the lowest and highest data within the 1.5 x IQR of the lower and 371 upper quartiles, respectively. Circles represent outliers. Linear mixed model with random intercepts 372 for canine individual and ductal segment was used. Pairwise comparison over all the individual 373 companions was implemented with Bonferroni's multiple comparisons correction. The level of 374 significant was defined as p < 0.05. Only statistically significant mean differences are indicated. 375

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Case nr.	Breed	Gender	Age at diagnosis
1	Skye Terrier	Intact female	9
2	German Shepherd Dog	Intact female	7
3	Mixed	Sterilized female	8
4	Jack Russel Terrier	Sterilized female	12
5	Bichon Frise	Intact female	11
6	Japanese Spitz	Sterilized female	6
7	Mixed	Intact female	9

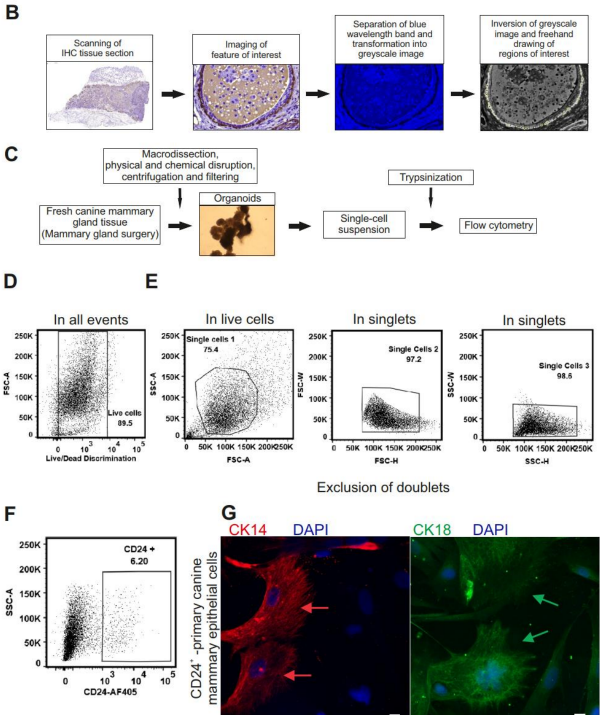


Figure S3: Determination of DAB intensity and isolation of epithelial cells for siRNA 379 experiments from canine mammary gland, Related to Figure 1 and Figure 2. A) Patient data of 380 canine cases selected for DAB intensity determination of basal myoepithelial markers α-SMA, CK5 381 and CK14. B) Scheme summarizing the protocol used to determine the DAB intensity (DABi) of 382 basal myoepithelial markers α-SMA, CK5 and CK14 from canine mammary gland tissue sections 383 stained with immunohistochemistry. Tissue section scans were generated with a Pannoramic 250 384 FLASH II digital scanner and features of interest imaged using Pannoramic Viewer. The 8-bit blue 385 wavelength band image was separated, transformed into greyscale and inverted using ImageJ, 386 followed by determination of the DABi from freehand-drawn cellular regions of interest (yellow in 387 image far right). C) Scheme summarizing the protocol used to isolate mammary epithelial cells from 388 freshly collected canine mammary gland tissue. **D-F)** FACS isolation strategy of CD24<sup>+</sup> canine 389 mammary epithelial cells showing the proportion of parent cells for each gate. D) Single-cellular 390 suspension of canine mammary epithelial cells gated for live cells using LIVE/DEAD® Fixable Near-391 IR Red. E) Doublets and larger cell clumps were discriminated in live cells population. Scatter Gate 392 393 display (left), FSC Gate display (middle) and SSC Gate display (right). F) Single-cells gated by CD24-AF405 for CD24<sup>+</sup> population representing epithelial cells. FSC-A = Forward Scatter Area; 394 395 SSC-A = Side Scatter Area; FSC-W = Forward Scatter Width; FSC-H = Forward Scatter Height; SSC-W = Side Scatter Width; SSC-H = Side Scatter Height. G) Immunofluorescence stainings of the 396 CD24<sup>+</sup> -primary canine mammary epithelial cells with specific antibodies against CK14 (red) and 397 CK18 (green). Nuclei were stained with DAPI (blue). Bar 10 µm. 398

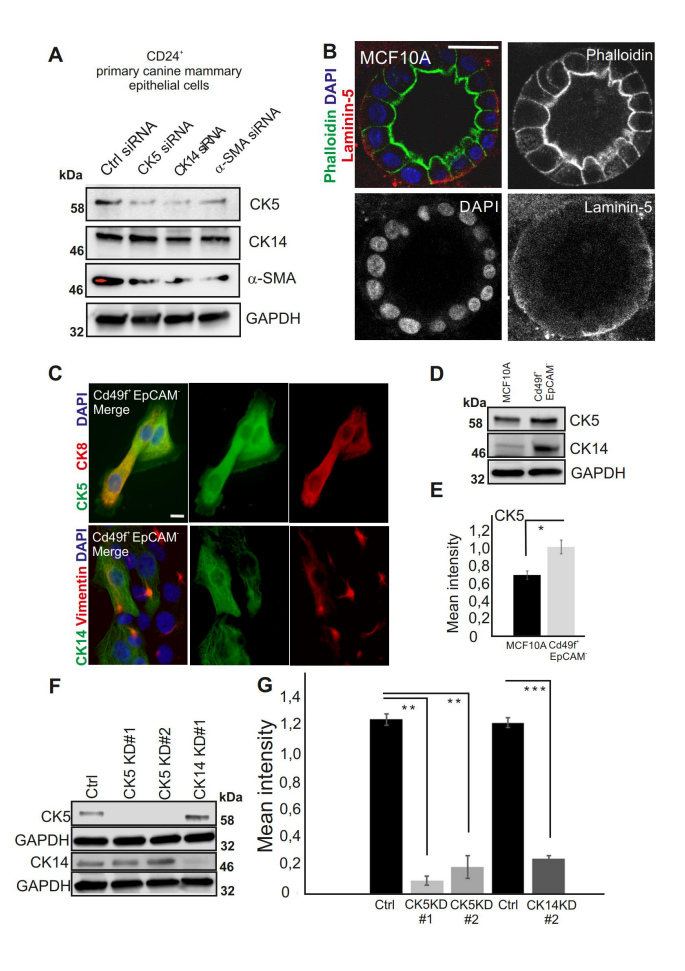
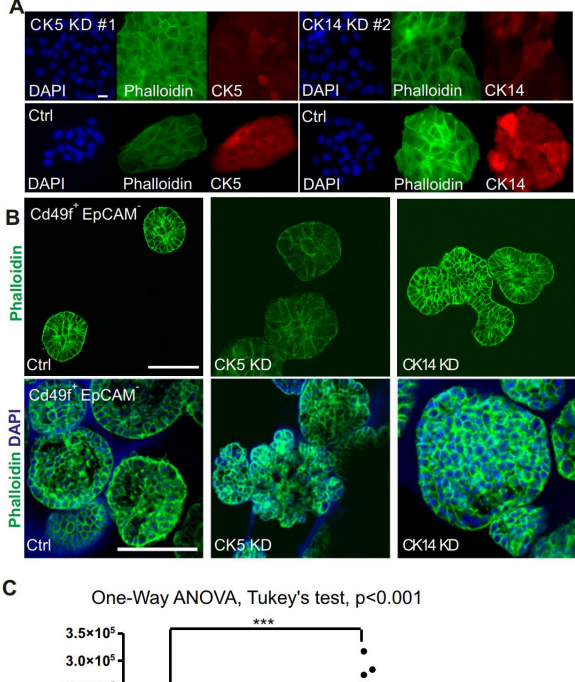


Figure S4: Knock-down of a-SMA, CK5 and CK14 from canine and human mammary 400 epithelial cells, Related to Figure 2. A) Depletion of primary canine mammary epithelial cells 401 (CD24<sup>+</sup>) by specific siRNAs against CK5, CK14 and  $\alpha$ -SMA on 2D cell culture conditions. Cellular 402 lysates were performed 4 days after application of siRNAs and protein levels were assessed in 403 Western blotting by specific antibodies. GAPDH acts as a loading control. B) MCF10A cells, cultured 404 for 2 weeks in 3D Matrigel, display hollow lumen and secretion of basement membrane proteins. 405 Specific antibody against laminin-5 was utilized for PFA-fixed spheroids and Phalloidin was used to 406 visualize actin cytoskeleton and DAPI nuclei. Bar 50 um. C) FACS-sorted CD49f<sup>+</sup> EpCAM<sup>-</sup> -407 mammary epithelial cell population enriched for basal progenitors, isolated from MCF10A cells and 408 stained with specific antibodies against CK5, CK8, CK14 and vimentin. Nuclei were visualized with 409 DAPI. Isolated cells display CK5/CK8 double positive cells and cells displaying either CK14 or 410 vimentin. Bar 20 µm. D) Comparison of parental cell line MCF10A and FACS-sorted CD49f<sup>+</sup> 411 EpCAM<sup>-</sup> cell population. Cellular lysates were utilized in Western blotting to assess the levels of 412 CK5 and CK14. GAPDH acts as a loading control. E) Quantification of CK5, related to Figure S3D. 413 Mean ( $\pm$  SEM) is shown; n=3; \*P<0.05 (paired t-test). F) Lentiviral-based RNA interference was 414 utilized to deplete CK5 and CK14 from CD49f<sup>+</sup> EpCAM<sup>-</sup> - mammary epithelial cell population 415 416 enriched for basal progenitors. Western blotting was used to confirm the efficiency of depletion. GAPDH was used as a loading control. G) Quantification of the CK5 and CK14 levels from ctrl 417 (CD49f<sup>+</sup> EpCAM<sup>-</sup>, enriched for basal progenitors), CK5 KD and CK14 KD cells, related to Figure 418 S3F. Mean (± SEM) is shown; n=3; \*\*P<0.01; \*\*\*P<0.001 (paired t-test). 419



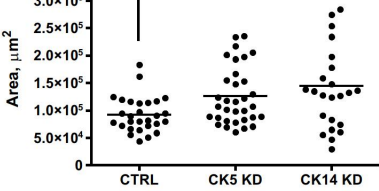


Figure S5: Depletion of CK5 and CK14 from CD49f<sup>+</sup> EpCAM<sup>-</sup> basal progenitor-enriched 422 MCF10A mammary epithelial cell population, Related to Figure 2. A) Immunofluorescence 423 stainings of the created knock down cell lines with specific antibodies against CK5 and CK14. Actin 424 cytoskeleton was visualized with Phalloidin and nuclei with DAPI. B) Depletion of CK5 and CK14 425 from 3D cultures of CD49f<sup>+</sup> EpCAM<sup>-</sup> basal progenitor-enriched MCF10A mammary epithelial cells. 426 Cultures were maintained for two weeks, after which they were fixed with PFA and stained with 427 Phalloidin (green) and DAPI (blue). Representative immunofluorescence images of two sets of 3D 428 mammospheres from separate experiments are shown. In the upper panel visualized with Phalloidin 429 and lower with Phalloidin and DAPI. Bar 100 µm. C) Measurement of spheroids area was performed 430 with Fiji-ImageJ 1.52p (National Institutes of Health, Bethesda, MD, USA, http://imagej.nih.gov/ij/, 431 1997-2019) software using free-hand selection tool and the built-in area measurement function. 432 \*\*\*P<0.001 (One-Way ANOVA analysis, Tukey's post-test. 433

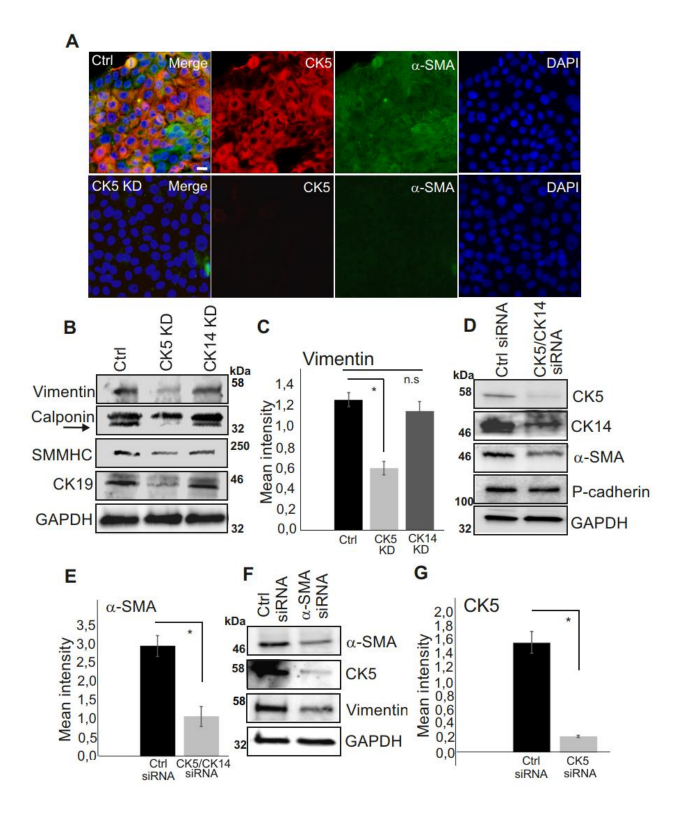


Figure S6: Depletion of CK5 leads to downregulation of markers for mature myoepithelium, Related to Figure 2 and Figure 3. A) Immunofluorescence stainings of ctrl (CD49f<sup>+</sup> EpCAM<sup>-</sup>, enriched for basal progenitors) and CK5 KD cells were stained with specific antibodies against CK5 and  $\alpha$ -SMA. Nuclei were visualized with DAPI. Bar 20 µm. B) Cellular lysates from ctrl (CD49f<sup>+</sup>

EpCAM<sup>-</sup>, enriched for basal progenitors), CK5 and CK14 KD cell lines were utilized in Western 439 blotting. Specific antibodies against vimentin, calponin 1, smooth muscle myosin heavy chain 440 (SMMHC) and CK19 were used in blotting. GAPDH was used as a loading control. C) Quantification 441 of vimentin levels, related to Fig. S5B. Mean (± SEM) is shown; n=3; \*P<0.05 (paired t-test); n.s.= 442 not significant. D) Scrambled siRNA and specific siRNAs against CK5 in combination with CK14 443 siRNA were utilized in MCF10A cells. Cellular lysates were performed four days after incubation 444 445 and protein levels for CK5, CK14, α-SMA and P-cadherin were detected in Western blotting with specific antibodies. GAPDH acts as a loading control. E) Quantification of the α-SMA levels showed 446 significant decrease upon CK5/CK14 depletion. Mean (± SEM) is shown; n=3; \*P<0.05 (paired t-447 test). F) Depletion of  $\alpha$ -SMA was performed with specific siRNAs in MCF10A cultures for four 448 days. Cellular lysates from control siRNA and  $\alpha$ -SMA siRNA- treated cells were used in Western 449 450 blotting and specific antibodies against  $\alpha$ -SMA, CK5 and vimentin were utilized to assess their levels. GAPDH was used as a loading control. Quantification of the CK5 levels showed significant decrease. 451 G) Quantification of CK5 levels, related to Figure S5F. Mean (± SEM) is shown; n=3; \*P<0.05 452 453 (paired t-test).

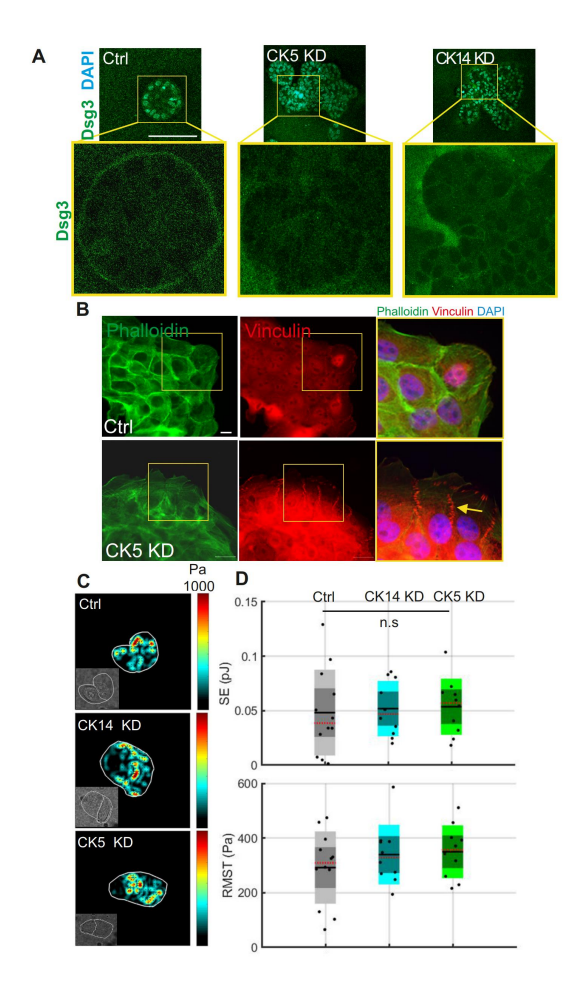


Figure S7: Knock down of CK5 impacts myoepithelial cell-cell junction proteins, Related to 455 Figure 4 and 5. A) 3D cultures of CD49f<sup>+</sup> EpCAM<sup>-</sup>/ basal progenitor-enriched MCF10A mammary 456 epithelial cells and CK5 and CK14 KD cells. Cultures were maintained for two weeks, after which 457 they were fixed with PFA and stained with Dsg3 (green) and DAPI (blue). Representative 458 immunofluorescence images of 3D mammospheroids are shown with magnifications from the 459 indicated areas below (yellow boxes). Bar 100 µm. B) Immunofluoresence microscopy was used to 460 detect cell-substrate adhesions from ctrl and CK5 KD cell lines. Specific antibody against focal 461 adhesion marker vinculin (red) was used. Phalloidin was used to stain actin cytoskeleton (green) and 462 DAPI (blue) for nuclei. Bar 20 µm. C) Traction force microscopy was applied to study cell-exerted 463 forces of the ctrl, CK5- and CK14 KD cells doublets. Representative force maps from ctrl, CK5- and 464 CK14 KD doublets after 3h incubation. D) Quantification of the Root Mean Square (RMS) tractions 465 and Strain Energy (SE). Data are presented as box-plots. n(ctrl)=12, n(CK5 KD)=10 and n(CK14 466 KD)=10. 467

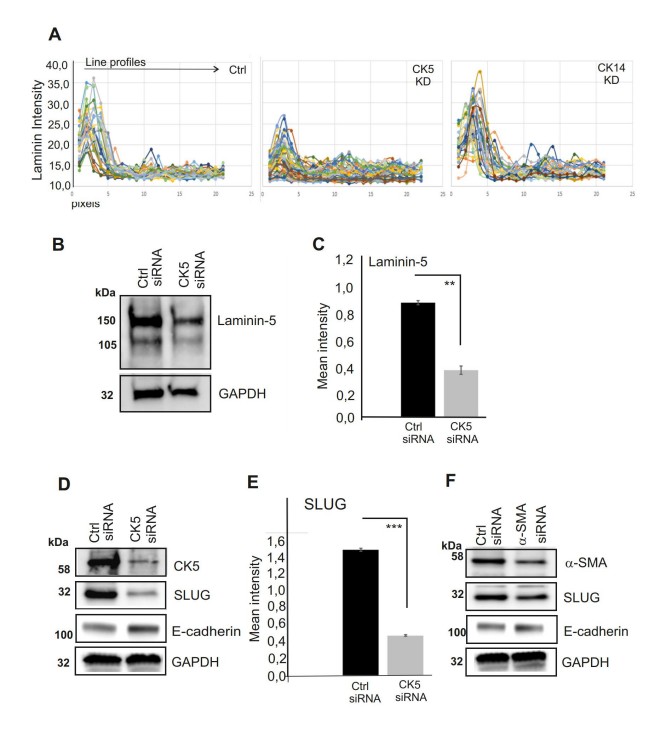




Figure S8: Loss of CK5 impacts expression of laminin-5 and SLUG, Related to Figure 6. A)
CD49f<sup>+</sup> EpCAM<sup>-</sup>/ basal progenitor-enriched MCF10A mammary epithelial cells, CK5 KD and CK14
KD cells were culture in 3D Matrigel for two weeks, after which they were fixed with PFA and
stained with laminin-5 (See also Fig. 6A and B). Lineprofiles were drawn from the edge of the
spheroid towards the center. 3-5 lineprofiles were drawn on each spheroid. Peak values on point 3
from lineprofiles were utilized for further analyses (Fig. 6B). B) Specific siRNAs against CK5 were

utilized in MCF10A cells. Cellular lysates were performed four days after incubation and specific 477 antibody against laminin-5 was used in Western blotting. GAPDH was used as a loading control. 478 Upper laminin-5 band (150 kDa) corresponds to the precursor form and lower band (105 kDa) to the 479 mature form. C) Quantification of the mature laminin-5 levels from CK5-depleted cells, related to 480 Figure S7B. Laminin-5 levels were divided with corresponding GAPDH values, and values of 481 untreated samples were set to 1. Mean (± SEM) is shown; n=4; \*\*P<0.01 (paired t-test). D) Specific 482 siRNAs against CK5 were utilized in MCF10A cells. Cellular lysates were performed four days after 483 incubation and specific antibodies against CK5, SLUG and E-cadherin were used in Western blotting. 484 GAPDH was used as a loading control. E) Quantification of the SLUG levels from CK5-depleted 485 cells, related to Figure S7D. SLUG levels were divided with corresponding GAPDH values, and 486 values of untreated samples were set to 1. Mean ( $\pm$  SEM) is shown; n=3; \*\*\*P<0.001 (paired t-test). 487 F) Depletion of  $\alpha$ -SMA was performed with specific siRNAs in MCF10A cultures for four days. 488 Cellular lysates from control siRNA and  $\alpha$ -SMA siRNA treated cells were used in Western blotting 489 and specific antibodies against  $\alpha$ -SMA, SLUG and E-cadherin were utilized to assess their levels. 490 GAPDH was used as a loading control. 491



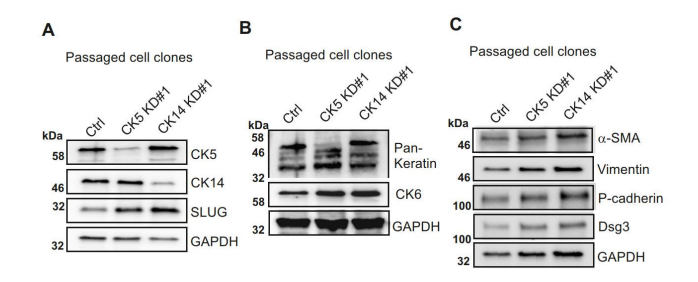


Figure S9. Passaged CK5 and CK14 KD cell lines undergo compensation by other cytokeratins,
Related to Figures 3-6. A) Cellular lysates from passaged cell clones of ctrl (CD49f<sup>+</sup> EpCAM<sup>-</sup>,
enriched for basal progenitors), CK5 KD and CK14 KD cells were utilized in Western blotting.
Specific antibodies against CK5, CK14 and SLUG were used to assess the levels of these proteins in
clones that had been in culture for several weeks. GAPDH was used as a loading control. B) Cellular

lysates from passaged cell clones of ctrl (CD49f<sup>+</sup> EpCAM<sup>-</sup>, enriched for basal progenitors), CK5 KD
and CK14 KD cells were utilized in Western blotting and Pan-keratin and CK6 antibodies were used
to detect possible compensation of the loss of CK5 and CK14. GAPDH was used as a loading control.
C) Cellular lysates from passaged cell clones of ctrl (CD49f<sup>+</sup> EpCAM<sup>-</sup>, enriched for basal
progenitors), CK5 KD and CK14 KD cells were utilized in Western blotting. Antibodies against αSMA, vimentin, P-cadherin and Dsg3 were used to assess the levels of these proteins in clones that
had been in culture for several weeks. GAPDH was used as a loading control.

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