Adhesion-growth factor crosstalk regulates AURKB activation and ERK signalling in re-adherent fibroblasts

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Aurora kinases despite their similarity have distinct roles in the cell cycle, which is regulated by cell-matrix adhesion and growth factors. This study reveals loss of adhesion and re-adhesion to differentially regulate Aurora kinases. AURKB activation that drops on the loss of adhesion recovers on re-adhesion in serum-deprived conditions but not in the presence of serum growth factors. A rapid 30 min serum treatment of serum-deprived cells blocks the adhesion-dependent recovery of AURKB, which negatively correlates with Erk activation. AZD mediated inhibition of AURKB in serum-deprived re-adherent cells promotes Erk activation and membrane ruffling, comparable to presence of serum. These studies thus define a novel adhesion-growth factor-dependent regulation of AURKB that controls adhesion-dependent Erk activation in re-adherent fibroblasts.

Keywords. Adhesion; AURKA; AURKB; Erk; growth factors

1. Introduction

Aurora kinase A, B, and C belong to an evolutionarily conserved family of serine/threonine kinases. Aurora kinases although sharing high structure similarity, perform distinct functions owing to their distinct subcellular localization in cells. The mitotic kinases Aurora Kinase B (AURKB) and Aurora Kinase A (AURKA) share about 71% identity in their catalytic domain (Carmena and Earnshaw 2003). Despite this similarity, AURKB and AURKA are found to be differentially activated in response to a variety of upstream stimuli and also regulate distinct downstream pathways during mitosis (Barr and Gergely 2007; Goldenson and Crispino 2015; Lan et al. 2004; Nair et al. 2009). AURKB is involved in chromosome condensation, sister chromatid cohesion, mitotic spindle assembly, regulation of spindle assembly checkpoint, cytokinesis, and daughter cell spreading (Crosio et al. 2002; Ferreira et al. 2013). AURKA is essential in centrosome maturation, centrosome separation, the formation of bipolar spindle assembly, and G2-M transition (Terada et al. 2003). All these functions are crucial for cell proliferation, with de-regulation of these kinases leading to incomplete and improper cell division leading to aneuploidy and affecting cell viability (Khan et al. 2011; Umstead et al. 2017).

Integrin-dependent signalling is an important mediator of cell-cycle progression during the G1 phase, in particular (Assoian and Schwartz 2001; Bae et al. 2014; Jones et al. 2019; Klein et al. 2009; Moreno-Layseca and Streuli 2014). Extracellular forces can regulate cell-cycle checkpoints through a focal adhesion kinase (FAK)/Rac signalling module (Bae et al. 2014) and the Hippo pathway on changing matrix stiffness (Dupont et al. 2011) (Aragona et al. 2013). Integrin-mediated adhesion is further seen to regulate cytokinesis and the re-spreading and repulsive migration of daughter cells (Aszodi et al. 2003; Hognas et al. 2012; Mathew et al. 2014; Pellinen et al. 2008; Taneja et al. 2016). While β1 integrin influences cytokinesis, αVβ5-positive reticular adhesions are essential for the normal progression of mitosis. Patterned 2D and 3D matrices reveal local tissue shape and mechanics to
regulate spatial differences in proliferation of cells (Aragona et al. 2013; Nelson et al. 2005, 2006). Exogenous stretching of epithelial cells is seen to stimulate cell cycle progression from G1 to S (Benham-Pyle et al. 2015) and G2 to M (Gudipaty et al. 2017). Cells are thus able to sense their surrounding ECM environment through integrin-associated adhesion complexes and determine whether to proceed through S phase or enter mitosis. Growth factors that activate RTKs can regulate integrin-mediated cell adhesion, spreading, and migration (Klemke et al. 1994; Mainiero et al. 1996; Trusolino et al. 1998). Conversely, integrins also support the activation of growth factor signalling that regulates cell cycle progression in fibroblasts (Assoian and Schwartz 2001; Danen and Yamada 2001; Gu et al. 1992). Integrin mediated cell cycle progression is mediated by the regulation of cyclin D1 expression through multiple pathways involving Erk, PI-3K, and the Rho family GTPases Rac, cdc42, and Rho (Danen and Yamada 2001; Roovers et al. 1999) and the negative regulation of p21cip1 and p27kip1 (Bottazzi et al. 1999).

During mitosis, human cells undergo mitotic cell rounding, decreasing their adhesion to extracellular matrix substrates (Li and Burridge 2019; Uroz et al. 2018). Spatiotemporal regulation of various mitotic kinase activity aids in the extensive cytoskeletal remodelling mechanisms that prevent detachment from epithelia, while aiding successful completion of cytokinesis (Champion et al. 2017; Petridou et al. 2019). AURKB is a dynamic mitotic kinase that has been reported to have multiple roles during different phases of the cell cycle. Apart from its extensively studied functions as a part of chromosome passenger complex (CPC), in early G1 phase AURKB associated with the cell cortex regulates cell spreading. It interacts and phosphorylates a formin, FHOD1, that is known to be essential to organize cytoskeleton after cell division (Floyd et al. 2013). Ferreira et al. report that, in late cytokinesis, the presence of a gradient of AURKB activity ensures that microtubules (MT) in the furrow region remain phosphorylated and the ones in the vicinity of extracellular matrix stay de-phosphorylated, restricting the MT growth at the cell-matrix interphase. This allows for the stabilization of focal adhesion, aiding co-ordinated daughter cells spreading (Ferreira et al. 2013). A membrane raft protein Flotillin-1 has been recently shown to regulate AURKB activity and CPC function providing a direct point of regulation between extracellular cues and cell cycle progression (Gomez et al. 2010). It however remains unclear if and how integrin-mediated cell-matrix adhesion can regulate Aurora Kinase B activation, and if the presence of serum growth factors can affect this regulation. Also of interest would be determining if these effects are dependent on the cell cycle.

In this study, we have hence tested if and how cell-matrix adhesion regulates AURKA/AURKB activity in anchorage-dependent WT-MEFs and if this contributes to integrin-dependent signalling (ERK, FAK, AKT) and cell spreading. The role serum growth factors have in regulating adhesion-dependent AURKB activation and the possible contribution cell cycle profile has on this regulation were also evaluated.

2. Materials and methods

2.1 Reagents

Human plasma fibronectin (Cat#F2006), nocodazole (Cat # M1404), DMSO (Cat # D2438), DAPI (Cat# D9542), Propidium iodide (Cat# P4170), and sparfloxacin (Cat# 56968) were purchased from Sigma, and Phalloidin-Alexa-488 (Cat# A12379) was from Molecular Probes (Invitrogen). Fluoromount-G used to mount cells for imaging was obtained from Southern Biotech (Cat# 0100–01). AZD1152 (AURKB inhibitor Cat# SML0268) was purchased from Sigma. RNAase-A (Cat# 9001–99–4) was purchased from USB Corporation. Immobilon Western blot substrate (Cat# WBKLS0500) was purchased from Millipore. Antibodies used for western blotting including anti-phospho-auro A (Thr288)/aurora B (Thr232)/aurora C (Thr198) (Cat #2914), anti-AURKB (Cat #3094), anti-phospho AKT—Ser473 (Cat # 9271) (1:2000 dilution), anti-phospho-FAK-Tyr397 (Cat# 3283), anti-phospho-p44/p42 ERK1/2 (Thr202/Tyr204) (Cat# 4370) (1:2000 dilution), anti-p44/p42 ERK1/2 (Cat # 4695) (1:2000 dilution), anti-FAK (Cat#3285) and anti-AKT (Cat# 4691) were purchased from Cell Signalling Technologies and used at 1:1000 dilution unless mentioned otherwise. Anti-AURKA (Cat #610939) was purchased from BD Transduction Laboratories used at 1:1000 dilution. Anti-beta actin (Cat# Ab3280) antibody was purchased from Abcam used at 1:2000 dilution. Secondary antibodies conjugated with HRP were purchased from Jackson Immunoresearch and were used at a dilution of 1:10000. Antibody used for immunofluorescence includes anti-phospho-p44/p42 ERK1/2 (Thr202/Tyr204) (Cat# 4370) from Cell signalling technologies used at 1:200 dilution. Secondary antibodies with Alexa conjugate (488 or 594) were purchased from Invitrogen Molecular Probes (Cat. No.
# A12379 and A12381) and were used at a dilution of 1:1000.

## 2.2 Cell culture

Wild-type Mouse embryonic fibroblasts (WT-MEFs) obtained from Dr. Richard Anderson (University of Texas Health Sciences Centre, Dallas TX), were cultured in DMEM (Cat# 11995-065) supplemented with 5% (v/v) fetal bovine serum (FBS) (Cat# 26140-079) and 1% (v/v) penicillin-streptomycin (Cat# 15140-122) (all from Invitrogen) at 37°C under 5% CO2 humidified atmosphere.

## 2.3 Suspension and re-plating of WTMEFs in serum starved or with serum conditions

For suspension assay, serum-starved (0.2% FBS) WT-MEFs (for at least 12 h) were detached with 1X trypsin-EDTA diluted with low serum medium and held in suspension for 30 minutes with 1% methylcellulose in low serum DMEM. Post incubation for respective time points cells were carefully washed twice with 0.2% FBS DMEM at 4°C to avoid clumping and collected. These washed cells were re-plated on dishes coated overnight with fibronectin at 4°C (10 μg/ml or 100 μg/ml as indicated in figure legends) for 15 min or 4 h. Cells were similarly grown and processed with serum (10% FBS) when required. When needed for western blotting these cells were lysed in the required volume of 1X laemmli, heated at 95°C for 5 min, and stored at −80°C.

## 2.4 Serum stimulation suspension assay

For serum stimulation suspension assay, serum-starved (0.2% FBS) WT-MEFs (for at least 12 h) were detached with 1X trypsin-EDTA diluted with low serum medium and held in suspension for 30 min with 1% methylcellulose in low serum DMEM. Post 30 min of suspension, a predetermined amount of 100% FBS was added to the suspension culture to adjust the final FBS concentration to 10%. Cells were suspended in 10% complete FBS or 10% heat-inactivated FBS (heat inactivation was done at 56°C for 30 min before the start of the experiment) media for additional 15 min to bring the total suspension time point to 45 min. Post 45 min (30 min low serum + 15 min 10% serum) incubation for respective time points cells were carefully washed twice with 10% FBS DMEM at 4°C to avoid clumping and collected. These washed cells were re-plated on dishes coated overnight with fibronectin at 4°C (10 μg/ml) for 15 minutes or 4 h. When needed for western blotting these cells were lysed in the required volume of 1X laemmli, heated at 95°C for 5 min, and stored at −80°C.

## 2.5 Cell cycle analysis by flow cytometry

For evaluating adhesion-dependent cell cycle profile, WT-MEFs grown in either 0.2% FBS DMEM or 10% FBS DMEM were detached using trypsin and held in suspension for 30 minutes. Post suspension cells were carefully washed, collected, and divided into three equal proportions: one processed immediately as suspension time point (SUS), one re-plated on 10 μg/ml fibronectin-coated dishes for 15 minutes (FN 15’), and last re-plated on 10 μg/ml fibronectin-coated dishes for 4 h (SA). These cells were washed twice with PBS and fixed using chilled 70% ethanol and stored at 4°C till further use (not more than 18 h). On the day of flow cytometer measurement cells were treated with 100 μg/ml RNAse A and labeled with 10 μg/ml of propidium iodide. DNA content was analyzed for cell cycle status in BD LSFRFortessa SORP cell analyzer. 10000 events were recorded for each treatment and time point to obtain the percentage of cells in different phases of the cell cycle. The cell cycle profiles were calculated by using ModFit software that gives percentages of cells in G0- G1, S, and G2-M phase and were compared across different treatments and time points. This method was used to evaluate the cell cycle profile of WT-MEFs that are stable adherent, held in suspension for 30 min, and re-adherent on fibronectin for 15 min in the presence and absence of serum growth factors.

## 2.6 Western blot detection of proteins

30 μl of lysate was resolved by 12.5% SDS-PAGE and transferred to PVDF membrane (Millipore). Blots were blocked with 5% milk in 0.1% Tween-20 containing Tris-buffered saline (TBST) for 1 hour at room temperature and incubated with required antibody diluted in 5% BSA at 4°C overnight. Blots were then washed thrice with TBST and incubated with anti-mouse HRP (AURKA) and anti-rabbit HRP (AURKB, Phospho-
AURK A/B/C, ERK, FAK, Phospho-ERK, Phospho-FAK) for an hour, followed by detection using chemiluminescent substrates from Millipore. LAS4000 (Fujufilm-GE) was used to image the blots and densitometric band analysis was done using Image-J software (NIH). Prism Graphpad software was then used to do the statistical analysis.

2.7 Immunofluorescence assay

WT-MEFs were serum-starved with 0.2% FBS containing DMEM (low serum DMEM) for 12 h, detached, held in suspension for 30 min in the absence or presence of AZD1152, and re-plated on fibronectin (2\mu g/ml) for 15 min. Re-adherent cells were fixed with 3.5% paraformaldehyde after 15 minutes of re-plating. Cells were permeabilized with PBS containing 5% BSA and 0.05% Triton-X-100 for 15 minutes and blocked with 5% BSA for 1 hour at room temperature followed by incubation with 1:200 rabbit anti-phospho-p44/p42 ERK1/2 (Thr202/Tyr204) antibody in 5% BSA for 3 h. Cells were finally stained with 1:1000 diluted secondary antibodies (anti-mouse Alexa-568) and 1:500 diluted phalloidin-Alexa 488 for 1 hour at room temperature. All incubations were done in a humidified chamber. Washes were done with 1X PBS at room temperature. Stained and washed coverslips were mounted with Fluoromount-G (Southern Biotech) and imaged using a Zeiss LSM 710 laser confocal-Anisotropy or LSM780 multiphoton microscope with a 63x objective.

2.8 Statistical analysis

All the analysis was done using Prism Graphpad analysis software. Statistical analysis of data was done using the two-tailed unpaired Student’s t-test for non-normalized data and when normalized to respective controls using the two-tailed single sample t-test. Distribution profile data was analysed using Chi-square test.

3. Results

3.1 Cell-matrix adhesion differentially regulates AURKB vs AURKA activation

To evaluate if cell-matrix adhesion can regulate Aurora Kinase A (AURKA) and Aurora Kinase B (AURKB) activity, serum-starved stable adherent (SA) wild-type mouse embryonic fibroblasts (WT-MEFs) were detached and held in suspension for 30 min (with 1% methylcellulose) (SUS 30'), and re-plated on fibronectin for 15 min (FN 15'). In the absence of serum growth factors, re-adhesion of cells is known to activate integrin-dependent downstream signalling. The effect that loss of adhesion and re-adhesion has on Aurora Kinase activity was tested by evaluating their phosphorylation at the Threonine 288 (AURKA) and Threonine 232 (AURKB) residues, known to be necessary for their activation (Ohashi et al. 2006; Yasui et al. 2004). In serum-starved WT-MEFs, AURKA activity drops (~60%) upon loss of adhesion and does not recover on re-adhesion (figure 1A), on the other hand, AURKB activity drops (~40%) upon loss of adhesion and is restored upon re-adhesion to fibronectin (figure 1B). AKT activation known to be regulated by adhesion expectedly drops on the loss of adhesion (SUS 30') and recovers on re-plating for 15 min (FN 15') (supplementary figure 1A). This suggests that integrin-dependent adhesion differentially regulates AURKA vs AURKB activity in WT-MEFs.

3.2 Adhesion-growth factor crosstalk regulates Aurora Kinase B activity

Knowing the overlap between adhesion and growth factor-mediated signalling, we tested the role serum growth factors could have on the adhesion-dependent regulation of Aurora kinases. On the loss of adhesion (suspension 30 min) activation of both AURKA and AURKB, seen to drop in serum-deprived conditions, (figure 1A and B) continue to decrease in the presence of serum growth factors (figure 1C and D). AURKA activity that does not recover on re-adhesion to fibronectin for 15 min, in the absence of serum, is also unaffected by serum growth factors (figure 1A and C). However, the recovery of AURKB activity on re-adhesion in serum-deprived conditions (figure 1B) is interestingly lost in the presence of serum growth factors (figure 1D). The AURKB activity in suspended cells without and with serum was comparable (supplementary figure 1C) and could hence not affect their recovery on re-adhesion. The drop in AKT activity on the loss of adhesion is also reduced in the presence of serum, while its recovery on re-adhesion is retained (supplementary figure 1B). Together they suggest integrin-mediated adhesion-dependent regulation of AURKB to indeed be affected by serum growth factors.
Figure 1. Adhesion-growth factor crosstalk-dependent regulation of AURKA and AURKB. Western blot detection (upper panel) and quantitation (lower panel) of (A and C) phosphorylation on Threonine 288 residues of AURKA (pAURKA), total AURKA, and (B and D) phosphorylation on the Threonine 232 residues of AURKB (pAURKB), total AURKB in the lysates from WT-MEFs stable adherent (SA), suspended for 30 min (SUS 30') and re-adherent on fibronectin for 15 min (FN 15') in presence of 0.2% (A and B) and 10% (C and D) FBS DMEM. The ratios of pAURKA/AURKA and pAURKB/AURKB are represented in the graph as mean ± SE from at least three independent experiments. Statistical analysis of all the above data was done using a two-tailed paired Student’s t-test and significance if any was represented in the graph (* p-value<0.05, ** p-value<0.01, *** p-value<0.0001).
3.3 Role of the cell cycle in adhesion-growth factor-mediated AURKB activation

Knowing the role of the cell cycle in regulation of Aurora kinases and the impact adhesion and growth factors could have on the same, we asked if and how their changing cell cycle profile could affect the observed regulatory crosstalk. Serum starvation of WT-MEFs, as expected, synchronize cells at the G1-S phase (Campisi et al. 1984; Griffin 1976; Langan and Chou 2011; Langan et al. 2017) with 76 ± 2.3 % of stable adherent cells in the G1-G0 phase and 3 ± 0.8 % in the G2-M phase (figure 2A). This profile does not change significantly when the cells are detached and held in suspension for 30 min, followed by re-plating on fibronectin for 15 min (figure 2A). The absence of any significant change in the cell cycle profile of fibroblasts suggests the regulation of Aurora Kinase B by adhesion is indeed independent of the cell cycle.

To further establish the differential regulation of AURKB we tested adhesion-dependent regulation of the cell cycle profile of WT-MEFs in the presence of serum. 50 ± 0.7 % of stable adherent WT-MEFs were seen to be in the G1-G0 phase and 43 ± 1.8 % in the S phase (figure 2B). This profile did not change significantly as these cells were held in suspension for 30 min, followed by re-plating on fibronectin for 15 min (figure 2B). In the presence of serum (figure 2B), the cell cycle profile was expectedly different as compared to that seen in the absence of serum (figure 2A).

The expression levels and activation of Aurora Kinases are seen to vary at different phases of the cell cycle (Carmena and Earnshaw 2003; Goldenson and Crispino 2015). A small but significant change in the cell cycle profile that the presence of serum causes could in part contribute to the differential effect serum growth factors have on adhesion-dependent AURKB activation. To evaluate this possibility, we tested if rapid (15min) stimulation of serum-deprived cells with serum growth factors, unlikely to affect their cell cycle profile, can affect the re-adhesion mediated regulation of AURKB. Serum-deprived (0.2% FBS DMEM) WT-MEFs suspended for 30 min were treated for 15min with 10% FBS and re-plated with 10% FBS DMEM on fibronectin for 15min. This treatment was found was enough to prevent the recovery of AURKB activity upon re-adhesion (figure 2C) as seen in WT-MEFs grown with 10% serum (figures 2C and 1D). This could have implications for serum growth factor-mediated fibroblast function. We hence tested if heat inactivation of serum (56°C, 30 min) can affect the adhesion-dependent AURKB activation (+ΔSer 30 min) and find that it does not (figure 2C). The AURKB activity in suspended cells without and with serum and on serum stimulation was comparable (supplementary figure 2A) and could hence not contribute to the recovery observed on re-adhesion. Focal adhesion kinase (FAK) activity is known to be regulated by integrin-mediated adhesion (Eliceiri 2001; Oktay et al. 1999) and was expectedly found to drop upon loss of adhesion comparably (supplementary figure 2B) and recover back upon re-plating on fibronectin irrespective of presence or absence of serum growth factors or heat-inactivated serum (figure 2D). This confirms serum growth factors to inhibit adhesion-dependent activation of AURKB, independent of the cell cycle.

3.4 Role of AURKB in regulating adhesion-growth factor-dependent ERK activation

To evaluate the functional relevance adhesion and growth factor-mediated regulation of AURKB activation could have in cells, we investigated if and how it could affect adhesion-dependent signalling or function. We tested the effect absence and presence of serum has on re-adhesion mediated activation of ERK and FAK (Eliceiri 2001) in WT-MEFs and the role AURKB activation could have in mediating the same. In the presence and absence of serum, FAK activation drops on the loss of adhesion comparably (supplementary figure 3C) and recovers on re-adhesion (figures 3C and 2D). Adhesion-dependent ERK activation in WT-MEFs was however seen to be differentially affected by the presence or absence of serum (figure 3A), like AURKB (figures 3B, 1B, and D). In low serum conditions ERK activation that drops on the loss of adhesion, further decreases on re-adhesion to fibronectin (figure 3A) as AURKB activation increases (figure 3B). Basal ERK activity was found to be significantly higher in WTMEFs in the presence of serum (figure 3A) but shows no significant change in its activation on the loss of adhesion or re-adhesion (figure 3A). In contrast, AURKB activation is seen to drop significantly in suspended cells and stays low on re-adhesion (figure 3B) in the presence of serum. This suggests the presence of a distinct inverse correlation between ERK and AURKB activation in re-adherent cells.

To confirm this, we evaluated the regulation of ERK on stimulation of serum-deprived cells with serum as well as heat-inactivated serum for 15 minutes. Both cause a drop in AURKB activity on re-plating as compared to serum-starved cells (figure 2C). This
Figure 2. Adhesion-growth factor dependent regulation of Cell cycle profile and AURKB activation. Representative histogram of 3 independent experiments and percentage of cells present in G2-M, S and G0-G1 phase are shown in table and graph from stable adherent (SA), suspended for 30 min (SUS 30’) and re-adherent on fibronectin for 15 min (FN 15’) in (A) serum starved and (B) with serum WT-MEFs. The graph represents the mean ± SE from at least 3 independent experiments. Statistical analysis was done using a two-tailed unpaired Student’s t-test and significance represented (* p-value<0.05, ** p-value<0.01). (C) Western blot detection (upper panel) and quantitation (lower panel) of phosphorylation on the Threonine 232 residues of AURKB (pAURKB), total AURKB and, (D) phosphorylation on Tyrosine 397 residues of FAK (pFAK) and total FAK in the lysates from: serum starved WT-MEFs (−Ser 12 h) suspended for 45 min and re-plated on fibronectin in presence of 0.2% serum, serum starved WT-MEFs suspended for 30 min followed by 15 min in heat inactivated 10% FBS DMEM (+Ser 30 min) and re-adherent on fibronectin for 15 min (FN 15’) in presence of heat inactivated 10% serum growth factors, serum starved WT-MEFs suspended for 30 min followed by 15 min in 10% FBS DMEM (+Ser 30 min) and re-adherent on fibronectin for 15 min (FN 15’) in presence of 10% serum growth factors or WT-MEFS suspended and re-plated in presence of 10% FBS DMEM. The ratios of pAURKB/AURKB and pFAK/FAK were normalized to respective SUS (equated to 1), and these values are represented in the graph as mean ± SE from four independent experiments. Statistical analysis of all the above data was done using the single sample t-test for normalized data and two-tailed unpaired Student’s t-test for non-normalized data and significance if any was represented in graph (* p-value<0.05).
regulation of AURKB is comparable to that seen in cells grown with serum (figure 3B). ERK activity in re-adherent cells is promoted in the presence of serum (figure 3D) further suggesting the adhesion-dependent AURKB activation to inversely regulate ERK activity in re-adherent cells. To further confirm the adhesion-dependent AURKB-ERK crosstalk, we tested the effect of AURKB inhibition on ERK activity and function. We used a specific AURKB small-molecule inhibitor AZD1152 (Wilkinson et al. 2007) to treat serum-deprived re-adherent cells (where AURKB is activated) (figure 1B) and compared their ERK activation to serum treated cells. AZD1152 treatment for 30 min inhibits AURKB (figure 4A and supplementary figure 4B), which causes a significant increase in ERK activity upon re-adhesion, comparable to serum treated cells (figure 4B). No significant effect of AURKB inhibition on ERK activation was seen in suspended cells (supplementary figure 4C) or on FAK activation in suspended as well as re-adherent WT-MEFs (supplementary figure 4A and 4D). Taken together, this confirms the presence of a unique AURKB-ERK regulatory crosstalk in re-adherent cells.

We further evaluated the localization of active phosphorylated ERK (pERK- Thr202/Tyr204) in re-adherent cells and tested the impact AZD1152 mediated AURKB inhibition and the resulting stimulation of ERK activation has on the same. Serum-deprived re-adherent cells make significantly fewer membrane ruffles unlike serum-treated WTMEFs (figure 4C and D). Active pERK is seen to prominently localize at membrane ruffles in serum treated cells where AURKB activation is blocked. In serum-deprived AZD1152 treated re-adherent cells, the inhibition of AURKB now causes cells to have a distinct ruffling phenotype with pERK actively recruited to membrane ruffles (figure 4C and D). Together these results confirm adhesion-dependent AURKB-ERK regulatory crosstalk to control ERK activity and localization downstream of integrin-mediated adhesion (figure 5).

4. Discussion

Aurora kinases are important regulators of cell division, single and collective cell migration (Barr and Gergely 2007; Carmena and Earnshaw 2003; Goldenson and Crispino 2015; Mahankali et al. 2015; Zhu et al. 2014) all of which are known to also be influenced by cell adhesion (Carstens et al. 1996; Jones et al. 2019; Li and Burridge 2019) and growth factor-mediated signalling (Golias et al. 2004; Jones and Kazlauskas 2001; O’Keefe and Pledger 1983; Rudland and Jimenez de Asua 1979; Schwartz 1997). The possible impact adhesion growth factor crosstalk, seen to regulate anchoradge-dependent signalling, has on AURK activation and function will have implications for multiple cellular pathways and processes. This study reveals AURKA and AURKB activity to drop on the loss of adhesion, AURK activation (but not AURKA) recovering on re-adhesion in serum-deprived conditions. This confirms AURKB to be regulated downstream of integrin mediated adhesion in mouse fibroblasts. This regulation is independent of the cell cycle, as serum-deprived fibroblasts showed no significant change in their cell cycle profile when suspended or re-adherent.

Knowing the effect adhesion growth factor crosstalk could have on cellular signalling, we tested and find the presence of serum did increase the percentage of S-phase cells, though the loss of adhesion and re-
Figure 4. Effect of AURKB inhibition on re-adherent ERK activity, localization, and membrane ruffling in WT-MEFs. Western blot detection (upper panel) and quantitation (lower panel) of (A) phosphorylation on the Threonine 232 residues of AURKB (pAURKB), total AURKB and, (B) phosphorylation on Threonine202/Tyrosine204 residues of p44/p42 ERK1/2 (pERK) and Total p44/p42 ERK1/2 (TotalERK) in the lysates from serum-starved WT-MEFs suspended for 30 min (with or without 2 μM AZD1152) and re-plated on fibronectin in presence of 0.2% serum (with or without 2 μM AZD1152) and WT-MEFS suspended and re-plated in presence of 10% FBS DMEM. The ratios of pERK/TotalERK and pAURKB/AURKB are represented in the graph as mean ± SE from three independent experiments. Statistical analysis of all the above data was done using the single sample t-test and significance if any was represented in graph (* p-value <0.05). (C) pERK detected using specific antibody against phosphorylation on Threonine202/Tyrosine204 residues of p44/p42 ERK1/2 (pERK) at membrane ruffles in re-adherent spreading WT-MEFs in presence of 0.2% FBS, 0.2% FBS + 2 μM AZD1152 or 10% FBS DMEM. (D) Percentage distribution of cells with ruffled and protrusion phenotypes in re-adherent WT-MEFs in presence of 0.2% FBS or 0.2% FBS + 2 μM AZD1152, or 10% FBS DMEM, was determined by counting 100 cells per time point from three independent experiments. Statistical analysis was done using Chi-square test two-tailed for distribution profile (D) data and p-values are as indicated (***p<0.001). Scale bar in (C) is set at 8 μm.
adhesion did not affect their cell cycle profile. In cells grown in the presence of serum re-adhesion mediated activation of AURKB is blocked. This when considered with the fact that basal activation of AURKB in suspended cells is unaffected by serum, suggests the serum-mediated regulation of AURKB to be unique to its crosstalk with integrin-mediated adhesion. The fact that a short 15min serum treatment can block the recovery of AURKB in serum-deprived re-adherent cells suggests this regulation to be independent of the cell cycle and mediated by a serum component. Heat inactivation of serum (56°C, 30 min) did not affect this regulation suggesting this component to be heat resistant. This could hence be a growth factor (like EGF, FGF, TGF) or small molecules like amino acids, sugars, lipids, or hormones (Honn et al. 1975). The dialysis or charcoal treatment of serum (Stoikos et al. 2008) could further help identify the serum fraction that carries this component.

Serum is seen to regulate Aurora Kinases, driving AURKA activation at the basal body of the cell ciliun in non-cycling G0/G1-phase mammalian cells, causing AURKA-dependent ciliary resorption (Pugacheva et al. 2007). Calcium-Calmodulin signalling is also seen to regulate the activation of AURKA (Plotnikova et al. 2010) and AURKB (Mallampalli et al. 2013), which downstream of integrins (Balasubramanian et al. 2007; Kwon et al. 2000; Shankar et al. 1993) through focal adhesion signalling (Giannone et al. 2004; Kirchoff et al. 1991; Naik and Naik 2003) could regulate AURKB. Cell-matrix adhesion could also regulate the spatial localization and activation of AURKB. Its localization at membrane protrusions could be mediated by formin (FHOD1) (Floyd et al. 2013) which is the major formin in mouse fibroblasts and seen to be active at sites of integrin engagement in cells (Iskratsch et al. 2013). Adhesion is also seen to regulate membrane raft trafficking which through resident proteins like Flotillin-1, can regulate AURKB (Gomez et al. 2010).

To establish the impact AURKB regulation could have on adhesion-dependent signalling in WTMEFs we tested the effect absence/presence of serum has on integrin-mediated ERK and FAK activation (Eliceiri 2001). The presence of serum promotes basal ERK activation significantly more than it affects FAK or AURKB activation. With serum regulating AURKB in re-adherent cells, ERK activation in re-adherent cells was seen to be inversely correlated to AURKB activation status. AZD1152 mediated inhibition of AURKB in serum-deprived cells confirms this crosstalk, by promoting ERK activation, that now localizes at membrane ruffles. This AURKB mediated regulation of ERK could be direct or indirect, which our ongoing studies are evaluating.

ERK, a serine/threonine kinase, is an important component of Ras-Raf-MEK-ERK signalling cascade that regulates cell proliferation, differentiation, migration, senescence, apoptosis and tumorigenesis (Chang and Karin 2001; Nishida and Gotoh 1993; Sun et al. 2015). The early adhesion-dependent activation of ERK is reported to specifically regulate lamellipodial protrusion in re-adherent cells. This is also seen to be associated with the detection of pERK in membrane ruffles of re-adherent cells (Mendoza et al. 2011). A pulsatile ERK activation is seen to be responsible for the dynamic generation of protrusions (Yang et al. 2018). Ras recruits the MAP kinase kinase kinase Raf, which phosphorylates and activates the MAP kinase kinase MEK, which then phosphorylates and activates ERK (McKay and Morrison 2007; Pullikuth and Catling 2007) at the plasma membrane. KSR1 (kinase suppressor of Ras 1) forms a ternary complex with B-Raf and MEK, that promotes activation of ERK and allows for its recruitment to the plasma membrane (Ray 2009) (McKay et al. 2009; Muller et al. 2001).

ERK regulates cell migration through its phosphorylation of myosin light chain kinase (MLCK), focal adhesion kinase (FAK), calpain, and p21-activated kinase (PAK1). ERK (ERK1 and ERK2) is activated by its phosphorylation on Thr<sup>202</sup>/185 and Tyr<sup>204</sup>/187 (human sequences) by MAPK/ERK Kinases (MEKs) (Minnigou and Blackwell 2020; Seger et al. 1992). Upon phosphorylation, ERK is either seen to homodimerize or heterodimerize (Khokhlatchev et al. 1998), which enhances ERK activity but reduces its translocation of ERK monomers to the nucleus. These phosphorylation
steps (Cirit et al. 2010; Kholodenko 2006), regulate the dissociation of MEK from ERK. This makes ERK very sensitive to the activation status of MEK. ERK inactivation is mediated by the MAPK phosphatases (MKPs) dependent removal of the Thr or Tyr phosphorylation’s (Caunt and Keyse 2013). ERK effectors localize in the cell cytoplasm, or the nucleus (Yoon and Seger 2006), with the nuclear translocation of ERK being vital in inducing gene expression and cell cycle entry. A regulatory crosstalk between AURKB-ERK1/2 downstream of adhesion could have implications in maintaining the fidelity of the cell cycle in anchorage-dependent cells and regulating adhesion-dependent cell spreading and/or migration. In silico sequence analysis of AURKB suggests that it could also be a possible substrate for ERK (unpublished data) as has been suggested in earlier studies (Eves et al. 2006). Ongoing studies in the lab are testing the same and its significance in cellular function.

During mitosis, Raf-1 is a key regulator of ERK activation (Galabova-Kovaes et al. 2006; Minden et al. 1994), through the Raf kinase inhibitory protein (RKIP) (Eves et al. 2006), also seen to regulate AURKB activation (Eves et al. 2006). ERK1/2 and AURKB are both individually involved in cell spreading (Ferreira et al. 2013; Fincham et al. 2000; Floyd et al. 2013; Xu et al. 2010) and migration (Matsubayashi et al. 2004; Sun et al. 2015; Xie and Meyskens 2013; Zhu et al. 2014) something their regulatory crosstalk could help fine-tune. Integrin engagement and downstream signalling pathways have been reported to be different in cells that are embedded in a 3D matrix (Davidenko et al. 2016; Duval et al. 2017; Kapaczynska et al. 2018; Martino et al. 2009). ERK1/2 activation in primary human fibroblasts grown in cell-derived 3D fibronectin was found to be significantly higher (Damianova et al. 2008). This combined with the fact that cell cycle progression is altered in cells embedded in 3D matrix (Desmaison et al. 2018; Moriarty and Stroka 2018) suggests that AURKB activity might be regulated differentially in 2D vs 3D microenvironments. Evaluating the AURKB-ERK1/2 crosstalk in 3D microenvironments could also be relevant for understanding the role it could have in maintaining the fidelity of the cell cycle in anchorage-dependent cells.

In anchorage-independent cancers, AURKB and ERK1/2 have been reported to synergistically enhance tumorigenic potential and aid in radio-resistance (Marampon et al. 2014; Niermann et al. 2011). In melanoma cells, BRAF/ERK axis has been shown to control AURKB expression at the transcriptional level (Bonet et al. 2012). In gynaecological cancers, MEK/ERK cascade has been shown to regulate AURKB signalling to sustain colony-forming potential, invasion, and migration along with altering their radiation response (Marampon et al. 2014). These reports further add to the possible role the AURKB-ERK1/2 crosstalk could have in diseases like cancer.

Much remains to be figured about the functional significance of this AURKB-ERK crosstalk in cells. This study in identifying their novel regulation downstream of adhesion and growth factors highlights a regulatory pathway that could have implications for multiple cellular processes.

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