Optimization of Formaldehyde Cross-Linking for Protein Interaction Analysis of Non-Tagged Integrin β1

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Formaldehyde cross-linking of protein complexes combined with immunoprecipitation and mass spectrometry analysis is a promising technique for analysing protein-protein interactions, including those of transient nature. Here we used integrin β1 as a model to describe the application of formaldehyde cross-linking in detail, particularly focusing on the optimal parameters for cross-linking, the detection of formaldehyde cross-linked complexes, the utility of antibodies, and the identification of binding partners. Integrin β1 was found in a high molecular weight complex after formaldehyde cross-linking. Eight different anti-integrin β1 antibodies were used for pull-down experiments and no loss in precipitation efficiency after cross-linking was observed. However, two of the antibodies could not precipitate the complex, probably due to hidden epitopes. Formaldehyde cross-linked complexes, precipitated from Jurkat cells or human platelets and analyzed by mass spectrometry, were found to be composed of integrin β1, α4 and α6 or β1, α6, α2, and α5, respectively.

1. Introduction

Protein-protein interactions are the basis for most cellular processes, including signalling, protein synthesis, and metabolism. Detailed knowledge of these protein networks is required in order to better understand diseases and develop adequate treatments. At present, protein-protein interactions are commonly investigated by yeast-two-hybrid approaches [1] and by in vitro binding studies [2]. However, these approaches are prone to false positive identifications because they do not take into account the temporal and local separations that occur in a living system. One tool to study protein-protein interactions in a physiological context is affinity enrichment of the protein of interest followed by detection of its binding partners using either immunodetection methods or mass spectrometry [3]. However, this classical immunoprecipitation method has two drawbacks. Weak interactions could be missed, if stringent wash conditions are applied. In contrast, nonstringent conditions may enable the identification of more proteins, but many of these could be false positives only binding the bait protein during sample preparation.

One approach to solve this problem is applying covalent cross-linking to intact cells and thereby stabilizing protein-protein interactions, including very weak and transient ones [3]. After this fixation step, highly stringent conditions can be used during cell lysis and affinity enrichment, minimizing the risk of identifying false positives. Several cross-linkers varying in spacer arm lengths, reaction groups, and other properties are commercially available. One of the shortest available cross-linkers is formaldehyde (2.3–2.7 Å), which has been used for a long time in histology and pathology to “freeze” the native state of tissues and cells [4]. The experimental conditions used in these applications lead to a very tight network of cross-links, which prevents the precipitation of one protein of interest as required for protein-protein interaction studies. However, lower formaldehyde concentrations (0.4–2% instead of 4%) and especially shorter reaction
times (minutes instead of hours) allow the utilization of formaldehyde as a cross-linker to analyze protein-protein interactions as shown by us and others [5–8].

The application of formaldehyde as a cross-linker has several advantages. Only closely associated proteins can be cross-linked due to the small size of formaldehyde. Furthermore, its high permeability towards cell membranes enables cross-linking in the intact cell, without addition of organic solvents such as dimethyl sulfoxide as necessary for other cross-linkers. Formaldehyde is also thought to allow very fast cross-linking and the stabilization of transient interactions [4]. Finally, formaldehyde is available in almost every laboratory at costs that amount to only a fraction of other cross-linkers. However, formaldehyde cross-linking is not yet an established standard method and many questions regarding the optimal experimental conditions and the usability of antibodies for pull-down of proteins after formaldehyde treatment remain. For example, epitopes recognized by antibodies raised against endogenous proteins after formaldehyde treatment remain. For example, conditions and the usability of antibodies for pull-down of and many questions regarding the optimal experimental cross-linking is not yet an established standard method necessary for other cross-linkers. Formaldehyde is also thought to allow very fast cross-linking and the stabilization of transient interactions [4].

In the present study, we report the optimization of a protocol applying formaldehyde cross-linking combined with immunoprecipitation and mass spectrometry (Figure 1(a)) to analyze the interaction network of integrin β1.

2. Materials and Methods

2.1. Cells and Reagents. Jurkat cells were grown in Dulbecco’s modified medium (GIBCO, high glucose) containing 10% fetal bovine serum (GIBCO), L-glutamine and penicillin/streptomycin. Human platelets were isolated from healthy human volunteers as described earlier [15]. This was approved by the University of British Columbia Research Ethics Board and informed consent was granted by the donors. Briefly, whole blood was drawn from the antecubital vein into 0.15% (v/v) acid-citrate-dextrose anticoagulant. Platelets were isolated by centrifugation and washed in physiological buffer. All anti-integrin β1 antibodies were monoclonal mouse anti-human antibodies provided by John Wilkins (Manitoba). Goat anti-mouse Alexa Fluor 680 was obtained from Molecular Probes and goat anti-mouse HRP was received from BioRad.

2.2. Modeling of Integrin β1 Structure. The integrin β1 structure was modeled on the structure of human integrin β3 [16] using SWISS MODEL [17, 18] and visualization was performed using the Swiss-Pdb Viewer Deep View (http://spdbv.vital-it.ch).

2.3. Formaldehyde Cross-Linking. Formaldehyde solution was obtained by dissolving 0.4% to 4% paraformaldehyde (Fisher Scientific) in PBS for 2 h at −80°C. The solution was filtered (0.22 μm), stored in the dark at RT and discarded after 4 weeks. For cross-linking, Jurkat cells were pelleted in a 50 ml reaction tube, resuspended in PBS and counted. Cells were centrifuged again and resuspended to 1 x 10⁷ cells/ml in formaldehyde solution. Cells were incubated with mild agitation for 7 min at RT and then pelleted at 1500 g and RT for 3 min, resulting in 10 minutes exposure to formaldehyde. The supernatant was removed and the reaction was quenched with 0.5 ml ice-cold 1.25 M glycine/PBS. Cells were transferred to a smaller tube, spun, washed once in 1.25 M glycine/PBS and lysed in 1 ml RIPA buffer (50 mM Tris HCl, pH 8.0, 150 mM sodium chloride, 1% NP40, 0.5% sodium deoxycholate, 0.1% SDS, 1 mM EDTA, protease inhibitors (Complete mini, EDTA-free, Roche Diagnostics)) per 1 x 10⁷ cells for 60 minute on ice. After 30 minutes, cell lysates were treated with 50 strokes using a Dounce homogenizer. Lysates were spun for 30 minutes at 20000 g and 4°C to remove insoluble debris. The supernatant was either used directly or stored at −80°C. Control cells
were treated exactly the same way, except that they were resuspended in PBS instead of formaldehyde solution. When platelets were used for cross-linking, ~1.5 × 10⁹ cells were resuspended in 10 ml formaldehyde solution and lysed in 1 ml RIPA buffer.

2.4. Immunoprecipitation, Western Blot Analysis and Silver Staining of SDS PAGE. Protein concentration was determined using a BCA assay (Pierce). The indicated amounts of antibodies and lysates were incubated for 1 h, after that 10 to 25 µl protein G agarose beads (Immobilized Protein G, Pierce) were added and immunoprecipitation was performed overnight. All steps were performed with mild agitation at 4°C. For mass spectrometric analysis, lysates were precleared by incubation with the same amount of beads for 2 h before the antibody was added. The supernatants of the immunoprecipitations were kept for analysis and the beads were washed either twice with PBS for western blot analysis or three times with RIPA buffer for mass spectrometric analysis. 4x reducing SDS Loading Dye (500 mM Tris HCl, pH 6.8, 8% SDS, 40% glycerine, 20% β-mercaptoethanol, 5 mg/ml bromophenol blue) was added to the beads as well as to the lysates and the supernatants and samples were incubated at either 65°C or 99°C for 5 min or 10–20 min, respectively, before they were separated by SDS PAGE using a 8% Laemmli gel [19]. For western blot analysis, proteins were transferred via a semi-dry procedure on polyvinylidene difluoride membranes (Pall Corporation), blocked for 1 h at RT with 5% milk powder in PBST (PBS, 0.1% Tween
20) and incubated with JB1A (0.1 µg/ml) overnight at 4°C. Membranes were either incubated with anti-mouse HRP or anti-mouse Alexa Fluor 680 (both 1:10000) for 1 h at RT. Membranes treated with anti-mouse Alexa Fluor 680 were scanned with a fluorescence scanner (Odyssey, LICOR) using excitation/emission wavelengths of 700 and 800 nm, whereas signals on anti-mouse HRP incubated membranes were detected with chemiluminescence solution (ECL, GE Healthcare). Quantitation of immunoblot analysis was performed using the software ImageJ [20]. Silver staining was performed using a modified protocol of Gharahdaghi et al. [21]. Briefly, gels were fixed with 10% acetic acid/10% methanol, washed with water and sensitized with 1 µg/ml dithiothreitol for 15 min. Gels were incubated in 0.1% (w/v) silver nitrate for 15 min and developed with 0.02% (w/v) paraformaldehyde in 3% (w/v) potassium carbonate until the desired staining had occurred. The reaction was stopped by addition of acetic acid.

2.5. In Gel Digestion and LC-MS/MS Analysis. Bands of interest in silver-stained gels were excised and destained as described [21]. Briefly, gel bands were incubated in a freshly prepared destaining solution (15 mM potassium ferricyanide/50 mM sodium thiosulfate) until the colour disappeared, washed several times with water and ammonium bicarbonate (pH 8.0) and chopped in smaller pieces. In-gel digestion was performed using standard procedures [22]. Briefly, samples were reduced with dithiothreitol, alkylated with iodoacetamide and digested with trypsin (Promega) in ammonium bicarbonate overnight. Peptides were extracted twice from the gel pieces, dried down in a vacuum centrifuge, reconstituted in 5% formic acid and STAGE-tip purified [23]. Separation and identification of peptides was performed by nano-HPLC MS/MS on an Agilent 1100 (Agilent, Santa Clara, CA) coupled to an FT-ICR (LTQ-FT, Thermo Electron Corporation, Waltham, MA) using a 15 cm long, 75 µm I.D. fused silica column packed with 3 µm particle size reverse phase (C18) beads (Dr. Maisch GmbH, Germany) with water:acetonitrile:formic acid as the mobile phase with gradient elution. Proteins were identified by extracting the Mascot generic format (MGF) files from the MS data using DTA Super Charge (part of the MSQuant open source project (http://msquant.sourceforge.net/)) and searching them against the ENSEMBL database with the X!Tandem algorithm embedded in the Global Proteome Machine (http://www.thegpm.org/). The protein expect score (log(e)) indicates the probability of false assignment of a protein: log(e) = −1 equates to a 1 in 10, log(2) = −2 to a 1 in 100 chance of a stochastic protein assignment. Known contaminants such as keratin were removed from the protein lists.

3. Results and Discussion

3.1. Formaldehyde Cross-Linking Conditions. The usage of formaldehyde as cross-linking reagent has to be evaluated in order to determine the optimal balance between highest yield of complex formation and lowest artefact generation [4]. Optimization has to be performed for each protein of interest, as its dependent on the physiological environment of the protein itself and can vary for example between cytosolic and membrane proteins. Three main parameters play a critical role during formaldehyde cross-linking: the reaction temperature, the incubation time and the formaldehyde concentration. The temperature dependency had been studied in our laboratory earlier and a difference between incubation at 37°C and 25°C could not be detected (unpublished results). Room temperature is advantageous, as it results in the most convenient and easiest approach possible. In addition, we chose not to increase the incubation time to more than 10 minutes, as the advantage of using formaldehyde as a cross-linker is the short reaction time it requires, which minimizes the formation of unspecific cross-links and allows the fixation of transient interactions. Moreover, model studies on peptides had shown that incubation time and formaldehyde concentration are complementary [24]. Therefore, we decided to limit our study to the usage of different concentrations of formaldehyde in terms of our model protein integrin β1 complex.

Jurkat cells were chosen for studying integrin β1 interactions, as this human T cell line expresses high amounts of this integrin and has been used extensively for its investigation before [25]. Different concentrations of formaldehyde (0.4% to 2%) were used to cross-link Jurkat cells. The lowest concentration was chosen as it had been shown earlier to result in the best protein loss/cross-linking yield balance [4], the highest was twice as high as formaldehyde concentrations shown to be successful earlier [5]. Cells were lysed under stringent conditions using RIPA buffer to destroy weak and noncovalent interactions, and protein amounts were determined. Lysates of formaldehyde treated cells contained lower amounts of protein than nontreated cells. This can be explained by the formation of insoluble complexes, for example, nuclear proteins being cross-linked to DNA, which were precipitated during lysis and removed in the insoluble pellet. This effect was visible during sample generation: lysis of nontreated cells using the stringent RIPA buffer led to the release of DNA, which formed a cloudy precipitate and could be easily removed. In contrast, a cloudy suspension was generated during lysis of formaldehyde treated cells and the pellet had a different consistency, which required a longer centrifugation period to become separated. Consistent with this observation, nuclear proteins were not detected in the lysate by immunoblot analysis. However, membrane proteins were overrepresented due to the loss of nuclear proteins (data not shown). We recommend using this difference in appearance as an early indication of successful cross-linking.

3.2. Detection of Formaldehyde Cross-Linked Integrin β1 Complexes. Optimization of the formaldehyde cross-linking protocol involving integrin β1 required a read-out of the cross-linking efficiency, for example, the detection of a complex containing the integrin. Formaldehyde cross-links are reversible during the standard sample preparation for SDS PAGE analysis (Figure 1(b)), which includes boiling in reducing Laemmli buffer [19], thus cross-linked complexes
would not be detected under these conditions [8]. However, by reducing the incubation temperature to 65°C, cross-linked complexes are not fully destroyed and remain detectable (Figure 1(c)) [5]. Membrane protein studies by gel electrophoresis are often performed at even lower temperatures (37–40°C). Initial immunoprecipitation experiments we had performed indicated that at 37°C the antibodies used for pull-downs would not dissociate into their heavy and light chains (data not shown). Instead, during gel electrophoresis the intact antibodies would migrate at molecular weights that would overlap with the cross-linked complexes and therefore interfere with their detection. Consequently, we decided not to use temperatures lower than 65°C in our experiments.

Jurkat cells treated with 2% formaldehyde were used to confirm the detection of cross-linked complexes. Lysates were incubated at 65°C and 99°C, respectively and analyzed by western blot using the antibody JB1A, which had been used for detection of integrin β1 in earlier studies [26]. We could recognize a higher molecular weight complex containing integrin β1 in samples treated for 5 min at 65°C, whereas this band was nearly undetectable after boiling for 10 minutes at 99°C (Figure 1(d)). Therefore, we concluded that we were visualizing a cross-linked complex containing integrin β1. However, integrin β1 was also found in the monomeric form at appr. 150 kDa after incubation at 65°C. This could be due to incomplete cross-linking, as the conditions applied during formaldehyde cross-linking do not lead to a high extent of protein cross-linking, leaving a large fraction of integrin β1 non-cross-linked. Alternatively, incubation at 65°C may lead to partial reversal of formaldehyde cross-links and release of integrin β1 even at a lower temperature.

3.3. Balance of Cross-Linking Efficiency and Protein Loss. Equal amount of lysates of the samples generated using different concentrations of formaldehyde were analyzed by immunoblotting (Figure 2(a)). No complex was detected in the control, in which cells were treated under cross-linking conditions but without formaldehyde. With 0.4% formaldehyde, a small amount of complex was detected, which increased with higher formaldehyde concentrations. At 0.8% formaldehyde, much more complex was visible. The amount increased slightly at higher concentrations of formaldehyde, but then settled, with no apparent difference between 1.2 and 2%. The signal intensities of monomeric integrin β1 and the complex were quantified, the values at 2% formaldehyde were set to 100% and relative intensities were calculated. These were plotted together with the total protein concentrations of the lysates to determine the optimal cross-linking parameters (Figure 2(b)). The amount of monomeric integrin β1 did not vary significantly between the different formaldehyde concentrations, even though the amount of the complex was increasing (Figure 2(b)). This apparent contradiction can be explained by the aforementioned observation that membrane proteins are enriched during formaldehyde treatment relative to other cellular components. Thus, by loading equal total protein amounts in each lane, increasingly higher amounts of total integrin β1 were applied. Unfortunately, this variation cannot be compensated using loading controls, as the exact amount of each cross-linked protein cannot be predicted. The decrease in protein concentration due to formaldehyde modification was most pronounced between 0% and 0.4% formaldehyde treatment, increased at 0.8% but stabilized at higher formaldehyde concentrations (Figure 2(b)). Comparing the loss of protein to the gain of integrin β1 complex (Figure 2(b)) implied that using a formaldehyde concentration between 1 and 2% should lead to the best cross-linking efficiency/protein loss balance, without major differences in this range. However, increasing the extent of cross-linking may also result in the formation of larger complexes by involving proteins that do not directly bind to integrin β1, including the cytoskeleton. This would lead to the formation of more extensive, heterogeneous

Figure 2: Optimization of formaldehyde concentration. Jurkat cells were treated with different concentrations of formaldehyde (PFA), lysed and the cell lysates were analyzed. (a) Anti-integrin β1 immunoblot of lysates (JB1A, anti-mouse HRP). (b) Western Blot results were quantified using ImageJ. Values obtained at 2% PFA were set to 100% and relative intensities were calculated. Protein concentration of the lysates had been determined using a BCA assay. Total protein concentrations and relative signal intensities of the western blot analysis were plotted in one graph to show the correlation between the parameters.
The use of a tagged, exogenous version of integrin can be used for interaction studies involving cross-linking. However, this was shown to be suitable for formaldehyde treatment, as studies utilizing myc-tagged proteins, the antibody 9E10 purification requires the antibody to still recognize its interaction partners using formaldehyde cross-linking and a 3.4. Analysis of Anti-Integrin β1 Antibodies

A closer look at the known properties of the anti-integrin β1 antibodies revealed that five of them had been shown to stimulate integrins when added to living cells, while two showed inhibitory character (Table 1). However, as we were planning to use these antibodies for precipitation of integrins from cross-linked and thereby fixed cells, these properties should not affect our experiments. More importantly, the epitopes of some of these antibodies are known and located at very different positions in the extracellular part of the protein: N29 and JB1B detect continuous peptide sequences close to the protein N-terminus (Figure 3(b)). In contrast, B3B11 and JB1B detect continuous sequences at the opposite end of the extracellular part of integrin β1 (Figure 3(b)), where also the discontinuous epitope of 2C18 is found [29]. B44 is known to recognize a different part of the protein than the other antibodies but the precise epitope is unknown. Since the B44 antibody is working in immunoblotting experiments, its epitope is expected to be composed of a continuous sequence [26]. In contrast, 3S3 cannot be used for immunoblotting indicating a discontinuous epitope.

We further modelled the integrin β1 structure on integrin β3 to gain insight into the three-dimensional location of the known epitopes (Figure 3(c)). As expected the epitopes are located in very different parts of the protein. Additionally it is known that the epitopes of N29 and B44 are hidden in the inactivated form of integrin β1 as on some cell lines they can only detect the receptor after it had become cross-linked complexes that would provide a larger surface to which nonspecific proteins can bind during sample processing. As a result, an increasing amount of abundant cytoskeletal proteins and common contaminants would be identified that would not be considered specific interactors. Therefore, to minimize such apparent artefact generation, we decided to perform the following investigations using more stringent conditions by applying 0.4% formaldehyde. This increases the likelihood of missed identifications of specific interactors of low abundance and of interactions of low stoichiometry. Even though higher formaldehyde concentrations would counter this effect, they would also result in more artefacts, hence differential analysis using multiple formaldehyde concentrations would still not be able to distinguish between specific interactors and artefacts. Instead, individual follow-up experiments for each identified protein would be required to determine its specificity. As the focus of this study was not to obtain an extensive list of putative interactors, but rather to demonstrate the general validity of the approach, we chose lower formaldehyde concentrations and high stringency. Users interested in maximizing the number of captured proteins should consider using higher formaldehyde concentrations instead.

### Table 1: Monoclonal anti-integrin β1 antibodies used in this study.

| Antibody | Activity | Epitope location | Reference |
|----------|----------|------------------|-----------|
| JB1B     | Stimulatory | Continuous (AA 671-705) | Wilkins et al. [26] |
| JB1A     | Inhibitory | Continuous (AA 82-87) | Ni and Wilkins [28] |
| B3B11    | Stimulatory | Continuous (AA 660-668) | Wilkins et al. [26] |
| N29      | Stimulatory | Continuous (AA 14-55) | Ni and Wilkins [29] |
| B44      | Stimulatory | Probably continuous/Diff. Region than N29 and JB1B/B3B11 | Wilkins et al. [26] |
| 6S6      | —        | —                | —         |
| 3S3      | Inhibitory | Probably discontinuous | Gao et al. [30] |
| 21C8     | Stimulatory | Discontinuous near JB1B/B3B11 | Wilkins et al. [26] |
activated [29]. To assess whether the antibodies would also be able to work after formaldehyde treatment, we had a closer look at the epitope sequences, which are known for four of them (Figure 3(b)). It had been shown that the amino acids, which are formaldehyde-modified first during the short incubation times used under cross-linking conditions, are lysine, cysteine and tryptophan [24]. The epitopes of N29 and JB1A are composed to a higher extent of these amino acids (20 and 30%), compared to 7% and 12% for B3B11 and JB1B, suggesting that N29 and JB1A are less likely to work after formaldehyde treatment.

3.5. Precipitation of Formaldehyde Generated Integrin β1 Containing Complex. Our next step was to determine whether the antibodies found to precipitate integrin β1 in the previous section can detect and pull down the integrin β1 complex. Jurkat cells were treated with or without 0.4% formaldehyde. Cells were lysed and the lysates were used for an immunoprecipitation using the antibody JB1B and a control pull-down which was performed without an antibody (Figure 4(a)). The higher molecular weight complex containing integrin β1 was only detected in formaldehyde treated cells and was only precipitated when the antibody was added. No complex was detected in the supernatant of the immunoprecipitation, indicating its complete pull-down. Furthermore, no loss in total integrin β1 precipitation was observed, indicating that formaldehyde treatment did not destroy the JB1B epitope. This had been expected, as the epitope does not contain many of the easily modifiable amino acids.

Figure 3: Analysis of anti-integrin β1 antibodies. (a) Test of antibodies for immunoprecipitation. 0.5 mg Jurkat cell lysate was incubated with 1 μg of the indicated antibody and a affinity purification was performed. Precipitations were analyzed by JB1A immunoblot (anti-mouse HRP). Two integrin β1 chains were detected in all immunoprecipitations. The heavy (HC) and light chains (LC) of the antibodies are indicated. (b) Amino acid sequence of the extracellular domain of human integrin β1. Known continuous epitopes are indicated. Underlined amino acids are known to become formaldehyde modified first. (c) Model of the extracellular part of human integrin β1, based on the crystal structure of Integrin β3 (Swiss Model). Indicated are the same epitopes as in (b).
As a next step, all of the other antibodies were used to precipitate integrin β1 from formaldehyde treated and non-treated Jurkat cells (Figure 4(b)). All of them were able to precipitate the target protein even after formaldehyde treatment, indicating that none of their epitopes was destroyed in such a way that it would prevent precipitation. Even for N29 and JB1A, whose epitopes contain a high percentage of easily modifiable amino acids, no fundamental loss in integrin β1 precipitation was observed. However, not all antibodies were equally able to precipitate integrin β1 containing complexes: B3B11, JB1A and 3S3 precipitated the complex completely, since no integrin β1 was detected in the high molecular range of the supernatant. Only partial precipitation of a high molecular weight complex was observed using the antibodies 2C18 and 6S6, where part of the complex was detected in the supernatant of the precipitation. No precipitated complex was detected using the antibodies N29 and B44. As these antibodies were able to precipitate integrin β1 of formaldehyde treated cells, it is unlikely that their epitopes were destroyed during the process. However, as mentioned earlier N29 and B44 have both been shown to detect activated integrin β1 on cells, whereas their epitopes are disguised on non-activated cells [29]. Stringent lysis of cells destroys this conformational limitation and allows detection by the antibodies. In contrast, cross-linking preserves the conformation and therefore a non-activated complex would...
be undetectable and not precipitated by these two antibodies. On the other hand, they could be able to precipitate an activated complex which in turn would not be recognized by some of the other monoclonal antibodies. This limitation of antibody usage does not only apply to formaldehyde cross-linking, but restricts every approach where protein complexes are precipitated by antibodies, including the traditional coimmunoprecipitation method which uses gentle lysis conditions to preserve protein interactions.

Finally we asked whether a higher degree of modification induced by higher concentrations of formaldehyde would impede the precipitation of integrin β1 using the antibody JB1B. Therefore we tested the ability of JB1B to pull down complexes from cell lysates, which were obtained after treatment of Jurkat cells with varying formaldehyde concentrations. JB1B was able to precipitate integrin β1 and the complex regardless of the formaldehyde concentrations applied, which was shown by analysis of the pull-downs as well as the supernatants of the immunoprecipitations (Figure 4(c)): no protein was detected in the supernatants, whereas integrin β1 was detected in the range of the monomeric protein and the high molecular weight complex in the immunoprecipitations. We concluded that JB1B is the most universal of the anti-integrin β1 antibodies we had tested, and therefore the most useful for formaldehyde cross-linking.

3.6. Mass Spectrometric Analysis of Complexes Containing Integrin β1. Western blot analysis had shown that we precipitate a high molecular weight complex containing integrin β1 after formaldehyde treatment. As the other components of this complex were unknown to us, we used mass spectrometry to identify them. The integrin β1 complex was precipitated by JB1B, separated by SDS PAGE and visualized by silver staining (Figure 5(a)). The indicated band was excised, in gel digested and analyzed by LC-MS/MS. Using Jurkat cells we performed this experiment in several independent repeats, and detected integrin β1 in every sample (Table S1). Furthermore we consistently identified integrin α4. The receptor formed by these integrins is VLA-4 (very late antigen 4), a known receptor found on T cells [31] and a high abundant integrin on Jurkat cells [25].

Two of these experiments also allowed the identification of additional proteins (Table 2). The listed proteins were not detected in control experiments using either nonformaldehyde treated cells or immunoprecipitations performed without the antibody when analysing gel bands at masses corresponding to the integrin β1 containing complex. Of these additionally identified proteins, integrin α6 is known to form a receptor termed VLA-6 together with integrin β1 on T cells, which has not been detected in Jurkat cells before [25]. Two additional integrin α subunits were detected only in dataset 2: integrin α5 (log(e) = −36.4) and α1 (log(e) = −1.7). Integrin α5 had been demonstrated to be expressed together with integrin β1 on Jurkat cells [25], whereas integrin α1 is known to form a collagen-recognizing receptor with integrin β1 on endothelial cells [32]. Even though only few peptides of each protein were sequenced, the identification of the integrin α subunits is reliable as they do not share high homologues sequences and all sequenced peptides were unique for each integrin α.

The other identified proteins are not known to interact directly with integrin β1 [13]. As a member of the cytoskeleton, actin is intertwined with integrin function and associated with integrins through adaptor proteins such as talin [10]. However, an adaptor was not consistently identified in these immunoprecipitations. Furthermore actin

**Figure 5: Analysis of integrin β1 complexes (CL) from Jurkat cells and human platelets.** (a) Jurkat cells were treated with or without 0.4% formaldehyde (PFA) before lysis. Immunoprecipitation was performed using 2 mg lysate and 5 μg of JB1B and analyzed by SDS PAGE. The indicated area was excised and analyzed by mass spectrometry. (b) Anti-integrin β1 immunoblot (JB1A, anti-mouse Alexa Fluor 680) of JB1B pull-downs performed on human platelets treated with and without 0.4% formaldehyde (PFA). 100 μg platelet lysate and 0.85 mg JB1B were used. Complexes containing integrin β1 are indicated by the arrow. (c) SDS PAGE analysis of JB1B immunoprecipitates from noncross-linked and cross-linked human platelets. Immunoprecipitation was performed using 0.7 mg lysate and 4.25 μg JB1B. The indicated band was excised and examined with mass spectrometry.
is a highly abundant protein and its identification has to be judged carefully. The same is true for GAPDH and heat shock protein HSP 90α, which are both abundant proteins and frequently found in immunoprecipitations. Thus we cannot exclude that these proteins are non-specific and frequently found in immunoprecipitations. Alternatively, proteins that do not colocalize but show medium to high affinity may form a non-cross-linked complex in the lysate that is sufficiently stable not to be destroyed by the incubation temperature of 65°C that is applied prior to gel separation. This scenario may also be true for most of the other proteins (Desmoglein-1, Filaggrin-2, Junction plakoglobin and C1orf68), which are highly expressed in skin cells. These could be contaminants similar to keratins (which were removed before data interpretation) occurring during sample preparation. The last identified protein arginase-1 is found in the liver and plays an important role in arginine metabolism [33], a connection to protein arginase-1 is found in the liver and plays an important role in arginine metabolism [33].

In addition to our studies using Jurkat cells, we wanted to apply our approach also to a different and physiologically more relevant system, and therefore used human platelets. Platelets express high amounts of integrins, which play an important role in platelet aggregation. Even though integrin β1 is not the most abundant integrin found on platelets, it forms three different receptors (α2β1, α5β1, α6β1) and is involved in platelet adhesion [34].

Human platelets were treated with and without formaldehyde and the antibody JB1B was used to precipitate integrin β1 from the lysates. In addition to the monomeric protein, a high molecular weight complex was detected in immunoblotting experiments, which was only visible after formaldehyde treatment (Figure 5(b)). We identified the contents of the complex precipitated by JB1B from human platelets by mass spectrometry (Figure 5(c), Table S2). Proteins identified in two independent experiments are listed in Table 3. Integrin β1 was detected in both pull-downs as expected. Furthermore, the integrins α6 and α2, which are known to form heterodimers with integrin β1 on platelets, were identified [34]. Thrombospondin-1, a known ligand of the integrin receptor α6β1, was also detected in the integrin β1 complex. As thrombospondin-1 is also the most abundant protein found in the α-granules of platelets [35], its identification may be merely due to its high abundance in the platelet lysate. Similarly, integrin α1β1 does not form a heterodimer with integrin β1, but together with integrin β3 forms glycoprotein IIb/IIIa, the most abundant platelet receptor. Integrin β3 was only detected in dataset 2 (log(e) = −40.2), however, the presence of both of these proteins could be solely due to their high presence in the platelet lysate as well. On the other hand we cannot exclude that integrin β1 interacts with one or both of these integrins forming multi-heterodimers.

### Table 2: Proteins identified in the integrin β1 complex precipitated from Jurkat cells. Formaldehyde cross-linked Jurkat cell lysate was used to enrich integrin β1 with JB1B. The integrin β1 complex was identified by SDS PAGE analysis and studied by LC-MS/MS and the GPM. Only proteins with a log(e) < −1 and which were identified in both datasets and not in control samples are listed.

| Protein                        | Accession       | Mr [kDa] | log(e) (Dataset 1) | log(e) (Dataset 2) |
|-------------------------------|-----------------|----------|--------------------|--------------------|
| Integrin β1                   | ITGB1 ENSP00000303351 | 88.4     | −39.8             | −118.3             |
| Actin, cytoplasmic 2          | ACTG1 ENSP00000331514 | 41.8     | −23.4             | −73.1              |
| Desmoglein-1                  | DSG1 ENSP00000257192 | 113.7    | −16.4             | −42.9              |
| Filaggrin-2                   | FLG2 ENSP00000373370 | 247.9    | −15.6             | −40.1              |
| Glyceraldehyde-3-phosphate dehydrogenase | GAPDH16 ENSP00000229239 | 36       | −7.4              | −18.2              |
| Arginase-1                    | ARG1 ENSP00000349446 | 35.6     | −7.3              | −15.1              |
| Junction plakoglobin          | JUP ENSP00000377507 | 81.7     | −6                | −7                 |
| Integrin α4                   | ITGA4 ENSP00000380227 | 114.8    | −5.1              | −6.3               |
| Heat shock protein HSP 90α    | HSP90AA2 ENSP00000216281 | 84.6    | −2.9              | −4.5               |
| C1orf68                       | C1orf68 ENSP00000354769 | 26.2     | −2.1              | −3.2               |
| Integrin α6                   | ITGA6 ENSP00000264106 | 126.5    | −1.6              | −2.4               |

Jurkat cells have been shown to express constitutively active integrin β1 [25]. However, it has to be considered that cell lines change their behaviour over time, thus it might be that the Jurkat cells used by them and by us differ in this aspect.
the experiments. Receptor VLA-5 with integrin β1 from human platelets. Using the transmembrane integrin β4 chain, we identified all known integrin α/β heterodimers on platelets. The identification of the adaptors talin-1 and kindlin-3 in one of the samples. However, these experiments were performed on platelets which were not intentionally stimulated, thus we do not expect high amounts of activated integrin β1 on these cells. Stimulation can nonetheless occur to a low degree during platelet preparation, which explains the detection of talin-1 and kindlin-3 in one of the samples.

4. Conclusion

Using the transmembrane integrin β1 complex as a model, our study addressed several critical factors in the formaldehyde cross-linking procedure and provides a practical guide to the optimization of such experiments.

Table 3: Proteins identified in the integrin β1 complex precipitated from human platelets. Integrin β1 was precipitated by JB1B from formaldehyde cross-linked platelets. The integrin β1 complexes were identified by SDS PAGE analysis and investigated by LC-MS/MS and the GPM. Only proteins with a log(e) < −2 and which were identified in both datasets are listed. *Two different isoforms were detected in the experiments.

| Protein                        | Accession   | Mr [kDa] | log(e) (Dataset 1) | log(e) (Dataset 2) |
|--------------------------------|-------------|----------|--------------------|--------------------|
| Thrombospondin-1               | THBS1 ENSP00000263056 | 129.3    | −46.2              | −140.1             |
| Integrin α6*                   | ITGA6 ENSP00000264106/ENSP00000364369 | 126.5/124 | −38.2              | −299               |
| Integrin β1                    | ITGB1 ENSP00000303351 | 88.4     | −29.8              | −201.1             |
| Immunoglobulin heavy chain C   | IGHV4-31 ENSP00000374990 | 43.9     | −9.6               | −10                |
| Hornerin                       | HRNR ENSP00000357791 | 282.2    | −9.4               | −20.2              |
| Integrin α2                    | ITGA2 ENSP00000296585 | 129.2    | −3.4               | −144.5             |
| Integrin α1βb                  | ITGA2B ENSP00000262407 | 113.3    | −2.8               | −22.8              |
| Filaggrin-2                    | FLG2 ENSP00000373370 | 247.9    | −4                 | −4.5               |

Table 4: Integrin β1 heterodimers detected in Jurkat cells and human platelets. *Only detected in one dataset.

| Detected in        | Integrin heterodimers | Ligands |
|--------------------|----------------------|---------|
| Jurkat cells       | β1α4 (VLA-4)         | Fibronectin, VCAM-1 |
|                    | β1α6 (VLA-6)         | Laminins |
|                    | β1α5 (VLA-5)*        | Fibronectin |
| Human platelets    | β1α6 (VLA-6)         | Laminins |
|                    | β1α2                 | Collagens |
|                    | β1α5 (VLA-5)*        | Fibronectin |

The identification of immunoglobulin heavy chain C is assumingly due to the immunoprecipitation, whereas hornerin and filaggrin-2 are both proteins expressed in the skin and could be contaminants caused during sample preparation. Three proteins which are known to interact with integrin β1 were only detected in dataset 2: integrin α5 (log(e) = −1.8), talin-1 (log(e) = −142.2) and kindlin-3 (log(e) = −18.5). Integrin α5 is known to form the receptor VLA-5 with integrin β1. With the detection of this α chain, we identified all known integrin β1 containing heterodimers on platelets. The identification of the adaptors talin-1 and kindlin-3 indicates again that we are able to precipitate integrin β1/adaptor protein complexes. However, these experiments were performed on platelets which were not intentionally stimulated, thus we do not expect high amounts of activated integrin β1 on these cells. Stimulation can nonetheless occur to a low degree during platelet preparation, which explains the detection of talin-1 and kindlin-3 in one of the samples.

By modifying SDS PAGE sample preparation conditions, we could detect an additional formaldehyde generated, high-molecular weight band containing integrin β1 by immunoblot analysis. LC-MS/MS analysis showed that this complex was mainly composed of integrins. Several integrin α subunits known to form heterodimers with integrin β1 could be detected in two different systems, Jurkat cells and human platelets (Table 4). The appearance of a broad band on the type of gel we had chosen did not provide any insights into heterogeneity and molecular weight of the corresponding integrin complex(es). Indeed, the band could have arisen from a single, multi-integrin complex containing all of the identified proteins, albeit the presence of several distinct complexes with similar molecular weight resulting in overlapping bands is more likely. Targeted experiments will be necessary to unequivocally answer this question.

One major focus of this study was to systematically test the applicability of monoclonal antibodies in precipitating formaldehyde treated endogenous, nontagged proteins. Two out of eight antibodies were expected to be less likely to work in precipitating integrin β1 from formaldehyde treated cells, since their epitopes contained amino acids that are known to be susceptible to formaldehyde modification [24]. However, we could not detect any difference in precipitation, indicating that destruction of epitopes is not an overwhelming problem in the formaldehyde cross-linking procedure at least for the panel of antibodies we had tested.

Two antibodies, which had been shown before to recognize only activated integrin β1 [29], did not pull-down the complex probably due to hidden epitopes in the complex. This finding, together with the observation that the analyzed complexes were mainly composed of integrin α and β chains, indicates that the majority of integrin β1 containing heterodimers found on Jurkat cells and human platelets are in an inactive state. However, in two experiments out of four we could identify talin-1, which is an adaptor protein that binds activated integrin β1. This suggests that formaldehyde cross-linking is a promising tool to study integrin activation and regulation. In future experiments Jurkat cell lines or platelets will be stimulated deliberately and the composition of the integrin β1 complexes will be analyzed. A quantitative
approach based on SILAC or iTRAQ labelling will then be applied to distinguish non-activated and activated integrin β1 and to allow timecourse studies.

By testing different formaldehyde concentrations for cross-linking, we realized that the balance between yield of cross-linked complex and the loss of protein was shifted in the case of integrin β1 compared to earlier studies: whereas 0.4% formaldehyde had been established as optimal concentration for studying myc-tagged soluble protein earlier [4], 1% formaldehyde seemed to be more suitable for integrin β1. While we used 0.4% formaldehyde for the current studies in order to minimize artefact generation, higher formaldehyde concentration may be used for future investigations. Indeed, varying the formaldehyde concentration is an excellent means not only to balance yield and protein loss, but also to distinguish direct and indirect interaction partners of a protein.

The multistep protocol for the optimization of formaldehyde cross-linking described here can be applied to any protein of interest. While the formaldehyde concentration is the most important variable that has to be optimized for each protein, detection of a formaldehyde cross-linked complex via western blot analysis simplifies this process. In addition, the application of antibodies for immunoprecipitations of modified proteins has to be monitored carefully, and the possibility that they do not recognize their antigen anymore has to be considered at all times. If these points are taken into account, however, formaldehyde cross-linking paired with immunoprecipitation will be a very powerful and reliable method to study protein-protein interactions, especially weak and transient ones, that can be established as a routine method in any laboratory.

**Abbreviations**

HRP: Horse radish peroxidase  
IP: Immunoprecipitation  
iTRAQ: Isobaric tag for relative and absolute quantitation  
LC-MS/MS: Liquid chromatography-tandem mass spectrometry  
Mr: Molecular weight  
PBS: Phosphate buffered saline  
PFA: Paraformaldehyde  
RIPA: Radio immunoprecipitation assay  
RT: Room temperature  
SDS PAGE: Sodium dodecylsulfate polyacrylamide gel electrophoresis  
SILAC: Stable isotope labelling with amino acids in cell culture  
SN: Supernatant  
VLA: Very late antigen.

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