Fc gamma receptor IIb modulates the molecular Grb2 interaction network in activated B cells

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1. Introduction

Development, survival and activation of B cells are tightly controlled by the B cell antigen receptor (BCR). The signals mediated by the BCR need to be precisely regulated to keep the balance between adaptive humoral immunity and tolerance. Hence, positive and negative regulatory effector proteins need to be accurately integrated into the signaling cascade. The BCR comprises an antigen-binding immunoglobulin subunit and an associated Ig-α/Ig-δ heterodimer that transduces signals via immune receptor tyrosine-based activation motifs (ITAMs). Engagement of the BCR leads to phosphorylation of ITAMs and subsequent activation of the spleen tyrosine kinase (Syk) [1]. A bona fide Syk substrate is the adapter protein Src homology 2 domain-containing lymphocyte protein of 65 kDa (SLP-65) [2] that upon phosphorylation provides docking sites for the SH2 domains of Bruton’s tyrosine kinase (Btk) and phospholipase C-γ2 (PLC-γ2) [3,4]. Formation of this trimolecular complex leads to mobilization of Ca2+ ions and activation of the NFκ-B and MAPK pathways [5].

BCR engagement can trigger opposed cellular fates like activation and differentiation or anergy and apoptosis, which depends on the developmental status of the B cell and concomitant engagement of other receptors [6]. Some of these receptors deliver co-activatory stimuli. CD19, for example, lowers the activation threshold of B cells by recruiting phosphoinositide-3-kinase (PI3K) [7,8]. PI3K catalyzes the production of phosphatidylinositol-3,4,5-trisphosphate (PIP3) at the cytosolic leaflet of the plasma membrane. PIP3 is a docking site for the Pleckstrin homology (PH) domains of Btk and PLC-γ2 and hence stabilizes their plasma membrane localization [9,10]. Limitation of BCR-induced activation is mediated by receptor proteins that harbor immunoreceptor tyrosine-based inhibition motifs (ITIMs). ITIMs recruit negative regulatory effector proteins via their SH2 domains [11]. For example, the sialic acid-binding immunoglobulin [Ig]-like lectin-2 (Siglec-2 or CD22) recruits the SH2 domain-containing protein tyrosine phosphatase-1 (SHP-1) which dephosphorylates several targets like Ig-α, Ig-δ and SLP-65 [12].

Another ITIM-containing receptor that has been extensively studied in the past is Fcγ receptor Iib (FcγRIib), which gets co-engaged if the BCR recognizes antigen that is already bound by IgG [13]. The phosphorylated FcγRIib ITIM recruits the SH2 domain-containing immunosuppressin 5-phosphatase (SHIP) which hydrolyzes PIP3 and hence destabilizes the plasma membrane localization of Ca2+-mobilizing enzymes [14,15]. This process is believed to prevent the production of new antibodies to a given antigen.
in a situation where class-switched and affinity-optimized antibodies are already available. Mice deficient for FcγRIIb mounted augmented humoral immune responses to both thymus-dependent and thymus-independent antigens [16]. Moreover, absence of FcγRIIb renders mice susceptible for autoimmune [17]. Interestingly, CD19, CD22 and FcγRIIb bind to the adapter protein growth factor receptor-bound protein 2 (Grb2) [18–20]. It has been shown that FcγRIIb-mediated SHIP recruitment is strongly compromised in Grb2-deficient DT40 cells [19]. However, the exact function of Grb2 in this context is not entirely understood.

Several studies elucidated the importance of Grb2 to regulate BCR-mediated signal transduction also independently of co-receptors. In complex with the adapter protein downstream of kinase-3 (Dok-3) Grb2 attenuates PLC-γ2-dependent production of inositol-1,4,5-trisphosphate [21]. Dok-3-deficient mice have increased serum IgM titers and enhanced humoral immune responses to T cell-independent antigens [22]. The regulatory function of Grb2 differs among developmental B cell stages. While in BCR-stimulated immature B cells Dok-3 appears to be the main Grb2-binding protein in mature B cells Grb2 is differently recruited by the transmembrane adapter protein linker for activation of T cells 2 (LAT2), which abolishes Grb2-mediated inhibition [23]. In memory B cells that express IgG- or IgE-containing BCRs Grb2 binds to the immunoglobulin tail tyrosine (ITT) motif which provides a co-stimulatory signal that appears to be essential for survival of these cells [24].

Recently, the Grb2 interactome in activated Bal17 B cells was elucidated by quantitative mass spectrometry analysis [25]. This study revealed that Grb2 interacts with a plethora of inhibitory and activatory signaling proteins. Notably a number of modulators of the PI3K signaling pathway were found among the 27 identified Grb2-associated proteins. These include positive regulators like the B cell adapter for PI3K (BCAP), Vav (seventh letter in the Hebrew alphabet) and the PI3K subunit p85 as well as negative regulators SHIP-1/-2 and SHP-2 are prominently bound to the pathway were found among the 27 identified Grb2-associated proteins. These include positive regulators like the B cell adapter for PI3K (BCAP), Vav (seventh letter in the Hebrew alphabet) and the PI3K subunit p85 as well as negative regulators SHIP-1/-2 and SHP-2 are prominently bound to the molecular Grb2 interaction network to inhibit BCR-mediated signaling.

In this study we elucidated the impact of FcγRIIb on Grb2-mediated signal integration in activated B cells by combining biochemical and quantitative mass spectrometry analysis. Our approach revealed that co-engagement of BCR and FcγRIIb augments the formation of ternary Dok-3/Grb2/SHIP protein complexes. Moreover, we could show by differential interactome analysis that the molecular Grb2 interaction network drastically differs in BCR- versus BCR/FcγRIIb-activated cells. Our data imply that Grb2 integrates more positive regulators of PI3K in BCR-stimulated cells whereas after co-engagement of FcγRIIb the negative regulators SHIP-1/-2 and SHP-2 are prominently bound to Grb2.

2. Material and methods

2.1. Cells, abs and reagents

Bal17.TR (kindly provided by Dr. Tony de Franco, University of California, San Francisco) were cultured in RPMI 1640 containing 10% FCS, 2 mM l-glutamine, 2 mM pyruvate, 50 μM B-ME, and antibiotics (cell culture reagents were purchased from Invitrogen). For SILAC experiments we used RPMI lacking arginine and lysine (Thermo Scientific). For retroviral infection of Bal17.TR cells we used a pMSCV-puro vector (Clontech) and constructs encoding mouse Grb2 [23] or the OneSTrEP-tagged variant [25]. BCR stimulation was performed for indicated times using 10 μg/ml F(ab)2 fragments of goat anti-mouse IgM or 15 μg/ml complete goat anti-mouse IgM (Jackson Immuno Research Laboratories).

Cells were lysed as described in [21]. For immunoprecipitation rabbit anti-Dok-3 S-20 (Santa Cruz Biotechnology) was used. For Western blot analysis we used anti-pYtr 4G10, anti-Grb2 3F2 (Millipore), anti-SHIP D1163, anti-Src Y418, Lyn Y509 (Cell Signaling Technology), and anti-Lyn 44 (Santa Cruz Biotechnology). For detection of mouse Dok-3 a polyclonal rabbit antiserum was kindly provided by Dr. Andre Veillette (Institut de recherches cliniques de Montréal). Recombinant Grb2-GST was prepared as described previously [26].

2.2. Mass spectrometric analysis

For stable isotope labeling of amino acids in cell culture (SILAC) Bal17. TR cells expressing a OneSTrEP-tagged Grb2 were cultured in RPMI 1640 medium devoid of arginine and lysine (Pierce) supplemented with 10% dialyzed FCS (Invitrogen), 1 mM pyruvate, and 4 mM glutamine. “Heavy” medium was supplemented with 12C6-D02N2 and 13C6-D02N4 arginine (Cambridge Isotopes) and “light” medium was prepared by adding equimolar levels of the corresponding non-labeled amino acids (Sigma-Aldrich). For affinity purification of OneSTrEP-tagged Grb2 106 cells from the respective cultures were stimulated with either goat anti-mouse F(ab)2 fragments to engage the BCR alone or with complete rabbit anti-mouse to co-engage BCR and FcγRIIb. Cleared cellular lysates were incubated with 200 μl of Strep-Tactin Superflow matrix (Iba BioTAgNology) for 1 h at 4 °C. After washing the immobilized Grb2 was eluted by incubating with 500 μl of D-desthiobiutin elution buffer (Iba BioTAgNology), pooled at a 1:1 ratio, and concentrated with ultrafiltration spin columns (Sartorius). Proteins were separated by one-dimensional SDS-PAGE (4–12% NuPAGE Bis–Tris Gel, Invitrogen) and each lane of the Coomassie-blue-stained gel was cut into 23 slices. All slices were reduced with 10 mM DTT for 55 min at 56 °C, alkylated with 55 mM IAA for 20 min at 26 °C and digested with modified trypsin (Promega) overnight at 37 °C. Following digestion peptides were injected into a C18 precolumn (1.5 cm, 360 μm o.d., 150 μm i.d., Reprosil–Porous 120 Å, 5 μm, C18-AQ, Dr. Maisch GmbH) at a flow rate of 10 μl/min. Bound peptides were eluted and then separated on a C18 capillary column (15 cm, 360 μm o.d., 75 μm i.d., Reprosil–Porous 120 Å, 5 μm, C18-AQ, Dr. Maisch GmbH) at a flow rate of 300 nl/min, with a gradient from 7.5% to 37.5% ACN in 0.1% formic acid for 60 min using an Agilent 1100 nano-flow LC system (Agilent Technologies) coupled to a LTQ-Orbitrap XL hybrid mass spectrometer (Thermo Electron). MS conditions were set as following: spray voltage, 1.8 kV; heated capillary temperature, 150 °C; normalized collision-induced dissociation (CID) collision energy 37.5% for MS/MS in LTQ. An activation q = 0.25 and activation time of 30 ms were used. The mass spectrometer was running in the data dependent mode. It automatically switched between MS and MS/MS acquisition. Survey MS spectra were acquired in the Orbitrap (m/z 350–1600) with the resolution set to 30,000 at m/z 400 and automatic gain control target at 5 × 105. The five most intense ions were sequentially isolated for CID MS/MS fragmentation and detection in the linear ion trap. Ions with single and unrecognized charge states were not included.

For quantification raw data were analyzed using the MaxQuant software (Version 1.0.12.31) in combination with Mascot search engine for peptide and protein identifications (Version 2.2.04, Matrix Science). IPI Mouse (Version 3.47) was used as a mus musculus sequence database. MS/ MS peak lists were filtered to contain at most six peaks per 100 Da intervals and searched against Mascot server. The MS mass tolerance was set to 7 ppm and MS/MS mass tolerance was set to 0.6 Da. Up to three missed cleavages of trypsin were allowed. Oxidized methionine and cysteine carbamidomethylation were searched as variable modifications. The modifications corresponding to arginine and lysine labeled with heavy stable isotopes could be treated as fixed modifications in the Mascot search, if applicable, after identification of SILAC pairs by MaxQuant. The false positive rate was set to 1% at the peptide level, the false discovery rate was set to 1% at the protein level and the minimum required peptide length was set to six amino acids.

2.3. Calcium measurements

106 cells were loaded in 700 μl RPMI containing 5% FCS, 1 μM Indo1-AM (Molecular Probes) and 0.015% Pluronic F127 (Molecular Probes) at 30 °C for 25 min. Subsequently, the cell suspension was diluted two-fold with RPMI 10% FCS and incubated for 10 min at 37 °C.
Cells were washed and prepared for measurements as described earlier [21]. Changes in the ratio of fluorescence intensities at 405 nm and 510 nm were monitored on a LSRII flow cytometer (Becton Dickinson) and analyzed with FlowJo (TriStar).

3. Results

3.1. Co-engagement of the BCR and FcγRIIb leads to increased formation of a ternary Dok-3/Grb2/SHIP complex

To examine the interdependence of FcγRIIb and Grb2 interactions we first analyzed the Dok-3/Grb2 association. Previous findings of our laboratory emphasize Dok-3 to be the main BCR-proximal Grb2-interacting protein under positive signaling conditions [21]. Moreover, Dok-3 and Grb2 bind to the FcγRIIb effector SHIP [27–29] implying that Dok-3/Grb2 might play a role under negative signaling conditions, too. To compare BCR versus BCR/FcγRIIb engagement we stimulated Bal17 B cells with two different protocols. To engage the BCR in the absence of FcγRIIb signals we used anti-IgM F(ab)2 that lacks the FcγRIIb-binding part. This procedure revealed very similar results like blocking the IgG binding site in FcγRIIb with anti-FcγR antibody (data not shown). For concomitant stimulation of BCR and FcγRIIb we treated cells with complete anti-IgM antibodies that bind to the FcγRIIb via its Fc part. We then prepared cellular lysates from these cells, immunopurified Dok-3, and subjected the samples to Western blot analysis. Probing the membrane with anti-Grb2 antibodies revealed an increased amount of co-purified Grb2 after BCR/FcγRIIb co-stimulation (Fig. 1A, first panel, lanes 5–7) compared to BCR engagement alone (lanes 2–4). This finding correlates with Dok-3 tyrosine phosphorylation (Fig. 1A, second panel). We looked for co-purified SHIP in our setup and anti-SHIP immunoprecipitation indeed revealed a stronger Dok-3/SHIP association in BCR/ FcγRIIb co-stimulated cells (Fig. 1A, third panel). Similar amounts of Dok-3 were confirmed as depicted in the fourth panel of Fig. 1A. To exclude that the increase in Dok-3 phosphorylation is merely due to augmented Lyn kinase activity, we assessed Lyn activation by Western blot analysis. We used phospho-specific antibodies that recognize the activating (Y397) and inhibitory (Y507) tyrosines, which are conserved in all Src kinases. This approach revealed no significant differences in Lyn activation in response to BCR crosslinking versus BCR/FcγRIIb co-crosslinking (Fig. 1B, first and second panels). Equal loading was confirmed by anti-actin immunoblotting (Fig. 1B, third panel). These results strongly imply that FcγRIIb utilizes Dok-3 by enhancing its tyrosine phosphorylation, which is not caused by a generally increased Lyn activity.

Our data show that the association of Dok-3 with both Grb2 and SHIP is augmented by co-crosslinking of BCR and FcγRIIb. Given the reported direct binding of Grb2 to SHIP this suggests formation of a ternary complex comprising Dok-3, SHIP, and Grb2. To test this, we made use of a Grb2-deficient B cell line (Bal17.TR) [30] that was either transfected with a Grb2 expression vector or empty vector as control. We performed again anti-Dok-3 immunopurifications and found that the amount of co-purified SHIP was strongly reduced in the absence of Grb2 expression (Fig. 2A, upper panel). This Grb2 dependency was most obvious when Dok-3 was immunopurified after co-engagement of BCR and FcγRIIb (lanes 5 and 6) or treatment with the phosphatase inhibitor pervanadate (lanes 7 and 8). The differences are neither due to varying amounts of purified Dok-3 (Fig. 2A, middle panel) nor to unequal SHIP expression (Fig. 2A, lower panel). Note that pervanadate treatment leads to additional posttranslational modifications of Dok-3, which is evident in slower migrating bands in the anti-Dok-3 immunoblot. To further confirm the importance of Grb2 for the Dok-3/SHIP association, we reconstituted Grb2 in vitro. Bal17 cells were treated with pervanadate to induce maximal tyrosine phosphorylation of SHIP and Dok-3. To cellular lysates recombinant GST-Grb2 or GST only as control was added. After immunoprecipitation of Dok-3 we detected co-purified SHIP by Western blot analysis. As shown in Fig. 2B, SHIP was only detected in the presence of GST-Grb2 (lane 3). These data show that Grb2 stabilizes the association of Dok-3 and SHIP leading to the formation of a ternary complex.

To test whether this finding has consequences for downstream signaling processes we monitored Ca²⁺ mobilization in the Bal17.TR transfectants described above. From Fig. 2C it is apparent that Grb2 attenuates Ca²⁺ flux after engagement of BCR alone (left graph). This is in line with our previous findings in DT40 and K46 cells [23]. However, the difference in Ca²⁺ profiles of Bal17.TR and reconstituted cells is significantly more pronounced after BCR/FcγRIIb stimulation (Fig. 2C, right graph). While BCR/FcγRIIb co-engagement compared to BCR stimulation already in absence of Grb2 expression limits the duration of

![Fig. 1. FcγRIIb co-engagement increases Dok-3 phosphorylation. (A) Bal17 B cells were left untreated (lane 1) or were stimulated via their BCR alone (lanes 2–4) or by co-engagement of BCR and FcγRIIb (lanes 5–7) for the indicated time points. As positive control cells were treated with pervanadate (lane 8). Dok-3 was immunopurified from cleared cellular lysates (CCL) and Western blots were tested for Grb2 (first panel), tyrosine phosphorylation (second panel), SHIP (third panel) and Dok-3 (fourth panel). (B) CCLs prepared from Bal17 cells that were either left untreated (lane 1) or stimulated for 3 min according to the protocol mentioned above (lanes 2 and 3) were subjected to Western blot analysis. Lyn tyrosine phosphorylation at position 397 (first panel) and position 507 (second panel) was analyzed by using phospho-specific anti-Lyn antibodies. As control the same samples were tested for actin (third panel).]
Ca\(^{2+}\) mobilization (compare black lines) in reconstituted cells concomitant engagement of BCR and Fc\(\gamma\)RIIb compromises the Ca\(^{2+}\) response drastically (compare gray lines). These results clearly confirm that Grb2 is also important for signal regulation under negative signaling conditions.

3.2. Fc\(\gamma\)RIIb co-ligation shifts the Grb2 interactome to negative regulatory proteins

Our results described above show that FcR\(\gamma\)RIIb co-engagement markedly increases formation of Dok-3/Grb2/SHIP complexes. Given the more than 20 Grb2-binding proteins that were recently identified in our laboratory\[25\] we now analyzed the impact of Fc\(\gamma\)RIIb on Grb2 interactions in more detail and employed differential interactome analysis by quantitative mass spectrometry. Briefly, we metabolically labeled Bal17.TR cells expressing OneSTrEP-tagged Grb2 in SILAC medium containing lysine and arginine with either "light" isotope (\(^{12}\text{C}_\text{6},^{14}\text{N}_\text{2}-\text{Lys}; ^{12}\text{C}_\text{6},^{14}\text{N}_\text{4}-\text{Arg}\)) or "heavy" isotope (\(^{2}\text{D}_\text{4},^{12}\text{C}_\text{6},^{14}\text{N}_\text{2}-\text{Lys}; ^{13}\text{C}_\text{6},^{14}\text{N}_\text{4}-\text{Arg}\)). The cells were then stimulated with either their BCR (light isotopes) or by BCR/Fc\(\gamma\)RIIb co-engagement (heavy isotopes) and Grb2 was purified from cellular lysates via a Streptactin™ matrix. Samples were pooled at a 1:1 ratio and after one-dimensional PAGE and trypsin digestion resulting peptides were identified by liquid chromatography-coupled tandem mass spectrometry (Fig. 3A). The metabolic labeling enables an unbiased relative quantification of co-purified proteins from differently treated cells as each identified peptide occurs with two distinct \(m/z\) (mass over charge) values. We were able to identify 26 Grb2-binding proteins (Table 1) that match the published Grb2 interactions\[25\].

Fig. 3. Quantitative mass spectrometric analysis of the Grb2 interactome in BCR- and BCR/Fc\(\gamma\)RIIb-stimulated cells. (A) Bal17.TR cells expressing OneSTrEP-tagged Grb2 were cultured in either "light" medium (\(^{12}\text{C}_\text{6},^{14}\text{N}_\text{2}-\text{Lys}; ^{12}\text{C}_\text{6},^{14}\text{N}_\text{4}-\text{Arg}\)) or "heavy" medium (\(^{2}\text{D}_\text{4},^{12}\text{C}_\text{6},^{14}\text{N}_\text{2}-\text{Lys}; ^{13}\text{C}_\text{6},^{14}\text{N}_\text{4}-\text{Arg}\)). Cells were then stimulated with either goat anti-mouse F(ab)\(_2\) fragments to engage the BCR alone or with complete goat anti-mouse to co-engage BCR and Fc\(\gamma\)RIIb. After lysis and OneSTrEP affinity purification samples were combined at a 1:1 ratio and proteins were separated by 1D-PAGE. Each gel lane was cut into 23 slices, which were digested and subsequently subjected to liquid chromatography-coupled tandem mass spectrometry (LC-MS/MS). The general analysis of the data obtained by the mass spectrometry experiments described in (A) is exemplary shown by representative sections of the spectra of "light" and "heavy" peptides derived from proteins Grb2 (B) and SHP-2 (C). The difference in the \(m/z\) values between the "light" and the "heavy" peptides are indicated within the spectra.
these interactors as exemplarily shown in Fig. 3B and C for a Grb2 and SHP-2 peptide, respectively. We then calculated the relative abundance of each corresponding pair of peptides (for detailed statistics see supplementary data). Heavy/light ratio values of >1 reflect increased and values of <1 reflect decreased amounts of co-purified proteins in the BCR/FcγRIIb-crosslinked sample compared to the BCR-stimulated sample. In Fig. 4, the mean ratios of two or three (*) independent experiments are illustrated as a bar plot. We identified three groups of proteins whose interaction with Grb2 is increased, decreased or not affected by combinatorial stimulation of BCR and FcγRIIb. The mass spectrometric approach confirmed our biochemical analysis described above as Dok-3 belongs to the group of proteins with augmented binding to Grb2 (ratio 1.48 ± 0.08). The increase of Grb2 association after FcγRIIb-co-engagement is more pronounced for the negative regulators SHP-1 (ratio 1.88 ± 0.21), SHP-2 (ratio 2.07 ± 0.14), and SHP-2 (ratio 1.68 ± 0.12), which fits the limiting role of FcγRIIb in BCR signaling. Consistently, the positive regulators P85α/β (ratio 0.65 ± 0.11) and Vav-1 (ratio 0.74 ± 0.12) belong to the group of proteins with FcγRIIb-dependent reduction within the Grb2 interactome. This also applies to PTP-PEST (ratio 0.76 ± 0.11) as well as the ubiquitin ligases Cbl-b (ratio 0.76 ± 0.16) and c-Cbl (ratio 0.69 ± 0.13). Notably Grb2 recruitment by other transmembrane proteins is affected by FcγRIIb co-engagement, too. We found diminished binding of Grb2 to CD19 (ratio 0.6 ± 0.10), CD72 (ratio 0.85 ± 0.04), and the receptor-type protein tyrosine phosphatase α (PTPRA) (ratio 0.71 ± 0.09). Moreover, Grb2 binding to putative regulators of the cytoskeleton dynamics and cellular trafficking like sorting nexin-18 (Snx-18) (ratio 0.64 ± 0.02), ASAP-1 (Arf-GAP containing SH3 domain, ankyrin repeats, and PH domain) (ratio 0.62 ± 0.03), and the RhOGAP ARHGAP-12 (ratio 0.76 ± 0.21) is decreased. The group of proteins that are purified from BCR- and FcγRIIb-stimulated cells in similar amounts include SLP-65, BCAP, HPK-1 (hematopoietic progenitor kinase-1), SHC, SHP-1, Gab-2-associated binder 2 (Gab-2), Dynamin-2/3, and SOS-1/2 (son of sevenless-1 and 2). Hence, our quantitative mass spectrometric approach shows for the first time a shift in the global interactions of a protein in response to co-crosslinking of an inhibitory co-receptor further indicating the importance of Grb2 for signal integration in B cells.

4. Discussion

The fundamental importance of FcγRIIb in keeping the balance between humoral immunity and autoimmune responses has mainly been attributed to the recruitment of SHIP. Our approach confirms the importance of SHIP and provides evidence for further alterations in the signaling cascade. We show that major parts of the molecular Grb2 interaction network are modified by FcγRIIb co-engagement which results in the differential integration of various regulators into the BCR signaling cascade. Consistent with these data the inhibitory function of FcγRIIb appears to require Grb2 as obvious from our Ca²⁺ flux analysis.

The Grb2-mediated signal integration starts with the SH2 domain-dependent rec localization of Grb2 to the plasma membrane. Despite a number of known interactors for the Grb2 SH2 domain, the mandatory Grb2-recruiting protein in BCR-stimulated cells appears to be Dok-3 because BCR-induced Grb2 relocalization hardly occurs in Dok-3-deficient DT40 B cells [21]. Our data show that BCR/FcγRIIb co-stimulation evokes a more pronounced tyrosine phosphorylation of Dok-3 than BCR stimulation alone. This is not related to an altered Lyn activity, which is consistent with earlier reports [31]. More likely FcγRIIb appears to control the Dok-3 microlocalization at the plasma membrane that increases the efficiency of its phosphorylation. A similar observation has been reported for Dok-1 that is also stronger phosphorylated after BCR/FcγRIIb stimulation leading to increased recruitment of the Ras GTPase-activating protein (RasGAP) and subsequent inhibition of the MAPK pathway [32]. From these data it appears that the most BCR-proximal PTK Lyn is utilized to mediate negative regulatory functions of FcγRIIb leading to augmented formation of Dok-1/RasGAP and Dok-3/Grb2 complexes. In the context of Grb2 this causes increased recruitment to the plasma membrane, which has fundamental consequences for further Grb2-mediated signal integration.

It has been reported earlier that the mouse ortholog of FcγRIIb itself harbors a binding site for the Grb2 SH2 domain and that SHP-1

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Table 1

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Fig. 4. FcγRIIb co-engagement changes the Grb2 interaction network. The quantitative mass spectrometry data of two or three (*) independent experiments were calculated as depicted in Fig. 3. The heavy/light ratios are normalized to Grb2 (ratio = 1) and error bars reflect the mean deviation. Values of > 1 reflect increased and values of < 1 reflect decreased amounts of co-purified proteins in the BCR/FcγRIIb-co-stimulated sample compared to the BCR-stimulated sample. The bold lines indicate the threshold for ratios that are significantly greater or smaller than 1.
might be lost. In this context it is noteworthy that the consensus Grb2 interaction with Fc of the mass spectrometry data those weakly co-purified proteins might be lost. In this context it is noteworthy that the consensus Grb2 binding site of mouse FcRlIib is not conserved among species leaving the role of this motif for SHIP-1 recruitment unclear. Our results imply that Grb2 stabilizes SHIP-1/FcRlIib complexes indirectly in alliance with Dok-3. We and others showed earlier that the Dok-3 PTB domain binds to SHIP-1 [21,28,33]. Consistent with these reports, which emphasize the importance of the Grb2 binding site in Dok-3 for SHIP-1 binding, we found Grb2 to be essential for this interaction. A similar ternary protein complex has been described for Shc, Grb2, and SHIP-1 [30]. However, FcRlIib co-engagement with the BCR appears to affect mainly Dok-3/Grb2/SHIP-1 complexes as we found Grb2 association with Shc to be unaltered. Therefore we propose that FcRlIib-dependent recruitment of SHIP-1 is a bidentated process. First, binding of the SHIP-1 SH2 domain to the phosphorylated FcRlIib ITIM and second, association with the plasma membrane resident Dok-3. Dok-3 and SHIP-1 are connected by Grb2 via its SH2 and C-terminal SH3 domain, respectively [19], leading to stable plasma membrane localization of SHIP. Consistent with this model we copurified markedly increased amounts of SHIP-1 with Dok-3 from BCR/FcRlIib co-stimulated cells. Our differential interactome analysis revealed a similar characteristic for the homologue SHIP-2. Indeed it has been proposed earlier, that SHIP-2 is also recruited to the FcRlIib ITIM [34]. However, the role of SHIP-2 in BCR or FcRlIib signaling remains elusive to date.

Our quantitative mass spectrometric analysis unveiled that FcRlIib co-engagement has considerable consequences for BCR-mediated signal integration beyond SHIP recruitment. Prominently upregulated in BCR/FcRlIib-co-stimulated cells is the SH2 domain-containing tyrosine phosphatase-2 (SHP-2). Unlike for its relative SHP-1 the exact role of SHP-2 in BCR signaling is unclear. Beside a proline rich motif interacting with the C-terminal Grb2 SH3 domain SHP-2 harbors a consensus binding motif for the SH2 domain that mediates adapter function leading to Ras activation downstream of a number of cytokine and growth factor receptors but neither the BCR nor FcRlIib [35–37].

Indications for a function of SHP-2 downstream of antigen receptors derive from studies in T cells [38]. T cells expressing a phosphatase-inactive SHP-2 variant develop normally but generate enhanced immune responses. It has been shown previously that FcRlIib co-engagement with the BCR leads to dephosphorylation of CD19 and Gab-2 [39,40]. An increased integration of SHP-2 might be the explanation for this observation. However, the exact role of SHP-2 in FcRlIib-mediated negative regulation requires further investigation.

Consistent with enhanced recruitment of the PI3K antagonist SHIP in BCR/FcRlIib-co-stimulated cells Grb2-mediated integration of the positive PI3K regulators p85 and Vav-1 is markedly reduced. Hence FcRlIib co-engagement appears to shift the whole PI3K regulation from activation to inhibition. Notably, Grb2 interaction with these regulators is SH3 domain-dependent as for SHP-2 and SHIP-1 [19,41–44]. This divergent signal integration of constitutively interacting proteins is probably based on the fact that stable multi protein complex formation often requires bidentated interactions. The FcRlIib-dependent SHIP recruitment discussed above is a very obvious example for this mechanism. It appears possible that integration of positive PI3K regulators is stabilized by Grb2 in context with the co-receptor CD19 in a similar way. CD19 is known bind the SH2 domains of Grb2 and p85 [18,45] and together with Vav and BCAP it orchestrates full PI3K activation [46]. CD19 tyrosine phosphorylation has been reported to be diminished after BCR/FcRlIib co-ligation [39]. Consistently, we found decreased binding of Grb2 to CD19 leading to reduced subsequent recruitment of PI3K activators and a reduction of these proteins within the Grb2 interactome. Hence, FcRlIib co-engagement appears to reduce the amount of PIP3 at the cytosolic side of the plasma membrane by both diminished PI3K activity and enhanced integration of SHIP (for summary, see Fig. 5).

Beyond marked alterations of PI3K regulation our analysis revealed FcRlIib-dependently decreased Grb2 binding to PTPTA, Snx-18, ARHGAP-12 and ASAP1. These proteins have not been studied in B cells so far and their putative function can only be deduced from studies in other cell types. Snx-18 and ASAP1 have been shown to regulate intracellular vesicle trafficking [47–49], PTPTA has been shown to regulate focal adhesion kinase (FAK) [50] whereas ARHGAP-12 is a putative modulator of Rho family GTPases [51]. Hence, it is tempting to speculate that these proteins are involved in regulation of BCR-mediated uptake of antigens as reorganization of the cytoskeleton and trafficking is important for this process [52]. Decreased Grb2-mediated integration of these proteins after FcRlIib co-engagement would be conclusive as antigen uptake is not required in this situation, but certainly this is a speculative interpretation.

Fig. 5. Model of Grb2-mediated signal integration in BCR- versus BCR/FcRlIib-stimulated cells. The dynamics of the molecular Grb2 network are summarized. Proteins that are more abundant in the Grb2 interactome of antigen-stimulated B cells are depicted in the left part of the model whereas the proteins that are more abundant after concomitant engagement of BCR and FcRlIib are shown on the right part of the model.
Collectively, our approach elucidates the dynamic characteristic of the molecular Grb2 network and emphasizes the complex impact of FcγRIIB co-engagement by shifting signal integration towards negative regulators. Based on these data we propose that differential plasma membrane relocation of Grb2 by Dok-3 or BCR co-receptors like CD19 is a crucial process to change integration of signal regulators. In conjunction with transmembrane or plasma membrane-associated proteins Grb2 appears to act as molecular glue to stabilize protein complexes. Here, binding of a Grb2 SH3 domain to a proline rich motif of a signal effector, which itself is bound to the same protein like the Grb2 SH2 domain increases the avidity of the whole protein complex. This could be the mechanism how functionally divergent co-receptors like FcγRIIB and CD19 utilize Grb2 to integrate either negative or positive regulators of BCR signaling with the ultimate aim to change the cellular response to BCR engagement.

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Appendix A. Supplementary data

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References