Phosphoproteomic analysis sheds light on intracellular signaling cascades triggered by Formyl-Peptide Receptor 2

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Formyl peptide receptors (FPRs) belong to the family of seven transmembrane Gi-protein coupled receptors (GPCR). FPR2 is considered the most promiscuous member of this family since it recognizes a wide variety of ligands. It plays a crucial role in several physio-pathological processes and different studies highlighted the correlation between its expression and the higher propensity to invasion and metastasis of some cancers. FPR2 stimulation by its synthetic agonist WKYMVm triggers multiple phosphorylations of intracellular signaling molecules, such as ERKs, PKC, PKB, p38MAPK, PI3K, PLC, and of non-signaling proteins, such as p47phox and p67phox which are involved in NADPH oxidase-dependent ROS generation. Biological effects of FPR2 stimulation include intracellular Ca2+ mobilization, cellular proliferation and migration, and wound healing. A systematic analysis of the phosphoproteome in FPR2-stimulated cells has not been yet reported. Herein, we describe a large-scale phosphoproteomic study in WKYMVm-stimulated CaLu-6 cells. By using high resolution MS/MS we identified 290 differentially phosphorylated proteins and 53 unique phosphopeptides mapping on 40 proteins. Phosphorylations on five selected phospho-proteins were further validated by western blotting, confirming their dependence on FPR2 stimulation. Interconnection between some of the signalling readout identified was also evaluated. Furthermore, we show that FPR2 stimulation with two anti-inflammatory agonists induces the phosphorylation of selected differentially phosphorylated proteins, suggesting their role in the resolution of inflammation. These data provide a promising resource for further studies on new signaling networks triggered by FPR2 and on novel molecular drug targets for human diseases.

Formyl peptides receptors (FPR1, FPR2, FPR3) are members of the GPCR family coupled to inhibitory G (Gi) proteins sensitive to pertussis toxin (PTX)1,2. “FPR2 binds efficiently the synthetic peptide” WKYMVm (W peptide) and is the most promiscuous member of the FPRs family, since peptides and lipids of different origin have been identified as FPR2 agonists3-5. FPR2 is highly expressed in cells of the immune system but it is also functionally expressed on cellular membranes of epithelial, neuronal and endothelial cells1. FPR2 was also detected onto nuclear membranes of anaplastic lung cancer CaLu-6 and human gastric adenocarcinoma AGS cells, where triggers intranuclear signaling cascades crucially involved in gene expression6. Furthermore, different studies highlighted the correlation between FPRs expression or function and inflammatory conditions, including inflammatory disorders, cancer, infections and other diseases7-9.

FPR2 stimulation induces multiple phosphorylations of several signalling proteins, which modulate critical intracellular functions such as cell growth, proliferation, intracellular communication, migration, differentiation, apoptosis, and survival5,10. For instance, the interaction of W peptide with its receptor elicits the activation of phospholipase A2, phospholipase C, and phospholipase D (PLA2, PLC, PLD)11,12, the stimulation of phosphatidylinositol-3-kinase (PI3-K)/protein kinase B (PKB or Akt) pathway13, and the activation of
protein kinase C (PKC), p38MAPK, extracellular response kinases 1 and 2 (ERK1/2) and c-Src. Some of FPR2-dependent phosphorylated molecules are non-signaling proteins, such as the cytosolic subunits p47phox and p67phox of NADPH oxidase complex whose phosphorylation requires FPR2-dependent activation of ERKs, PKCo, and PKCδ. Activated NADPH oxidase complex generates reactive oxygen species (ROS), which play a crucial role in tyrosine kinase receptors (TKRs) transactivation in several cell lines. In CaLu-6 cell line, FPR2 stimulation by W peptide induces Epidermal Growth Factor Receptor (EGFR), p47phox and c-Src phosphorylation. Phosphotyrosine residues of EGFR provide docking sites for recruitment, phosphorylation, and triggering of Signal Transducer and Activator of Transcription 3 (STAT3) transcriptional factor. In human prostate epithelial cell line PNT1A, FPR2 activation induces the phosphorylation of Y1313, Y1349 and Y1356 residues of Hepatocyte Growth Factor Receptor (c-Met) and triggers STAT3, PLC-γ1/PKCα and PI3K/Akt pathways. Furthermore, N-fMLP stimulation of FPR1 induces the phosphorylation of cytosolic Y951, Y966, and Y1175 residues of Vascular Endothelial Growth Factor Receptor 2 (VEGFR2), which constitute the anchoring sites for signaling molecules. These, in turn, activate PI3K/Akt and PLC-γ1/PKC intracellular pathways.

Phosphorylation represents the most frequent reversible covalent modification of proteins observed in eukaryotic cells. About 2–5% of the human genome codes for kinases and phosphatases and about 30% of the total content of proteins in eukaryotic cells is phosphorylated at any given time. However, in eukaryotic cells the overall level of tyrosine and threonine phosphorylated residues in proteins is very low compared to the level of phosphoserine.

Characterization of the phosphorylation status of cells plays a key role in the study of intracellular signaling cascades, in the understanding of the molecular mechanisms responsible of human disorders and, in turn, for the appropriate pharmacological approach. Although intracellular signaling pathways triggered by W peptide in FPR2-expressing cells are widely investigated, a complete analysis of the phosphoproteome is not currently available. Herein, we apply a phosphoproteomics approach to investigate phosphorylated proteins in WKYMVm-stimulated CaLu-6 cells. We identified 290 phosphoproteins and 53 unique phosphopeptides mapping on 40 proteins and we prove that selective phosphorylation on Ser82 of Heat Shock Protein-27 (HSP-27), Ser139 of Mini-Chromosome Maintenance protein 2 (MCM2), Ser339 of Oxidative-Stress-Responsive kinase 1 (OSR1), Ser608 of Retinoblastoma (Rb) and Ser170 of Myristoylated Alanyl-Rich C-Kinase Substrate (MARCKS), specifically depends on FPR2 stimulation by WKYMVm. The interconnection between some of the signalling readouts identified was demonstrated by using specific signalling inhibitors. Furthermore, we show that FPR2 stimulation with Annexin A1 (ANXA1) or Lipoxin A4 (LXA4), two anti-inflammatory FPR2 agonists, induces the phosphorylation of the selected differentially phosphorylated proteins.

Our overall data demonstrate that network-based global phosphoproteomic analysis can contribute to the identification of new intracellular pathways elicited by FPR2 which, in turn, could lead to the identification of new targeting strategies for FPR2-dependent human diseases.

Methods

Cell culture and reagents. CaLu-6 cell line was obtained from American Type Culture Collection (ATCC, Manassas, VA, USA). Cells were grown in Dulbecco's modified Eagle's medium (DMEM) (Thermo Fisher Scientific, Monza, Italy) supplemented with 10% foetal bovine serum (FBS) (Invitrogen). Once reached 80% confluence, cells were serum starved for 24 hours, and stimulated for 5 minutes with WKYMVm (Primm, Milan, Italy) at the final concentration of 10 μM, or with Lipoxin A4 (Santa Cruz, CA, USA) at the final concentration of 1 μM or with Annexin A1 (Abcam) at the final concentration of 10 nM. In other experiments, serum-deprived cells were preincubated for 16 hours with PTX (Sigma) at a final concentration of 100 ng/mL, or with 10 μM WRW4 (Sigma) (Primm, Milan, Italy) for 15 minutes, or with 50 μM PD98059 (Sigma) for 90 minutes, or with 5 Rottlerin μM (Sigma) for 1 hour before the stimulation with W peptide. SDS-PAGE reagents were purchased from Bio-Rad (Hercules, CA, USA). Anti-phosphoHSP-27 (S28), anti-phosphoMCM2 (S139) and anti-phosphoRb (S608) were from Cell Signalling Technology (Denvers, MA, USA). Anti-phosphoOSR1 (S339) was purchased from Signalway Antibody (Baltimore, MD, USA) and anti-phosphoMARCKS (S170) were from GeneTex (Irvine, CA, USA). Anti-tubulin, anti-cyclin D, anti-cyclin E, anti-rabbit and anti-mouse antibodies were purchased from Santa Cruz Biotechnology (Santa Cruz, CA, USA). Protein A–horseradish peroxidase was from Thermo Scientific (Little Chalfont, Buckinghamshire, UK).

Phosphoproteins enrichment. The enrichment of phosphorylated proteins was performed by a phosphoprotein purification kit (Qiagen, Hilden, Germany), according to the manufacturer's instructions. Briefly, whole lysates were obtained from 24-hours serum starved CaLu-6 cells stimulated or not with W peptide for 5 minutes. Cellular proteins were incubated with the appropriate amount of Phosphoprotein Lysis Buffer for 30 minutes at 4 °C and gently vortexed every 10 minutes. Lysates were centrifuged at 10000xg for 30 minutes at 4 °C and protein concentration was determined. Lysates (2.5 mg) were loaded onto Phosphoprotein Purification Column to allow the binding of phosphorylated proteins and the column was washed with the appropriate buffer as described in the manufacturer's instructions. Finally, phosphoproteins were eluted by adding to the column 500 μL of Phosphoprotein Elution Buffer. This step was repeated four times and the concentration of the four enriched phosphoprotein fractions was determined. To concentrate samples, an ultrafiltration step was performed by centrifuging the eluted fractions, at 10000xg for 10 minutes.
Alixouts of enriched phosphoprotein samples (~50 μg) from 24-hours serum starved Calu-6 cells, treated or not with W peptide for 5 minutes, were precipitated by adding pre-chilled acetone (six volumes) for 16 h at –20 °C. Following centrifugation for 10 min at 8,000 × g at 4 °C, protein pellets were resuspended in 100 μL of 50 mM NH4HCO3 pH 8.2. Samples were then subjected to tryptic digestion with 10 mM DTT (1 h at 55 °C) and alkylation with 7.5 mM iodoacetamide (20 min at room temperature in the dark). Enzymatic hydrolyses were performed on reduced and alkylated samples by adding TPCK-treated trypsin with an enzyme-substrate (E/S) ratio of 1:100 (w/w) for 3 h and 1:50 for 16 h at 37 °C.

**High-resolution nanoLC-Tandem mass spectrometry.** Mass spectrometry analyses of tryptic digests (2 μg) were performed as previously described25 on a Q-Exactive Orbitrap mass spectrometer equipped with an EASY-Spray nanoelectrospray ion source (Thermo Fisher Scientific, Bremen, Germany) and coupled to a ThermoScientific Dionex UltiMate 3000RS LC nano system (Thermo Fisher Scientific).

**Data processing.** The acquired raw files were analysed with Proteome Discoverer 2.1 software (Thermo Fisher Scientific) using the SEQUEST HT search engine as described25. The HCD MS/MS spectra were searched against the Homo sapiens database available in UniprotKB (20,413 reviewed entries, release 2018_01, 31-Jan-2018) assuming trypsin (Full) as digestion enzyme with two allowed number of missed cleavage sites and a minimum peptide length of six residues. The mass tolerances were set to 10 ppm and 0.02 Da for precursor and fragment ions, respectively. Carbamidomethylation (+57.021464 Da) of cysteine was set as static modification. Phosphorylation of serine, threonine, and tyrosine (+96.980 Da) and acetylation of lysine (+42.011 Da) were set as dynamic modifications. False discovery rates (FDRs) for peptide spectral matches (PSMs) were calculated and filtered using the Target Decoy PSM Validator Node in Proteome Discoverer with the following settings: maximum Delta Cn of 0.05, a strict target FDR of 0.01, a relaxed target FDR of 0.05 and validation based on q-value. Localization and best site probability for variable post-translational modifications within peptides were performed with the ptmRS tool, integrated in Proteome Discoverer26. The Protein FDR Validator Node in Proteome Discoverer was used to classify protein identifications based on q-value. Proteins with a q-value of 0.01 were classified as high confidence identifications and proteins with a q-value of 0.01–0.05 were classified as medium confidence identifications. The resulting protein table was then filtered based on the presence of phosphorylated peptides. Phosphopeptide changes following WKYMVm treatment were only considered when modifications were identified in at least two out of three replicate injections in treated sample and not identified in replicate injections of untreated sample.

**Bioinformatic analysis.** Interaction network analysis of proteins identified by LC-MS/MS were performed by the FunRich open access software (http://funrich.org/index.html). Molecular functional enrichment of identified proteins based on gene ontology categories was performed by using the Network analyst software (https://www.networkanalyst.ca)27.

**Western blotting.** Whole lysates were obtained as previously described28. Calu-6 cells were serum-deprived for 24 hours and then stimulated with 10 μM W peptide for 5 minutes. In other experiments, cells were pretreated with the appropriate amount of specific inhibitors before the stimulation with WKYMVm. Briefly, cells were washed in cold phosphate buffered saline (PBS) and lysed by incubation with RIPA buffer (50 mM Tris–HCl, pH 7.4, 150 mM NaCl, 1% NP-40, 1 mM EDTA, 0.25% sodium deoxycholate, 1 mM NaF, 10 μM Na3VO4, 1 mM phenylmethylsulfonylfluoride, 10 μg/ml aprotinin, 10 μg/ml pepstatin, 10 μg/ml leupeptin) for 45 min at 4 °C29. Bio-Rad protein assay was used to determine proteins concentration (BioRAD, Hercules, CA, USA). Fifty micrograms of whole lysates were resolved on 10% SDS-PAGE and proteins were transferred by blotting onto PVDF membranes. Membranes were incubated with phosphospecific primary antibodies and probed with appropriate horseradish peroxidase-conjugated secondary antibodies. Proteins were detected by autoradiography and bands densitometry was estimated using a Discover Pharmacia scanner equipped with a sun spark classic densitometric workstation30. The same filters were reprobed with an anti-tubulin antibody to normalize the amount of loaded proteins.

**Statistical analysis.** All the data are representative of three or more independent experiments and are expressed as means ± Standard Error Mean (SEM). Statistical analyses were evaluated by one-way analysis of variance (ANOVA) performed with GraphPad Prism 7 (GraphPad Software, San Diego, CA, USA). A p value of less than 0.05 was considered statistically significant.

**Results**

**Phosphoproteomic analysis by high-resolution nanoLC—MS/MS.** The global phospho-signaling response triggered by WKYMVm in Calu-6 cells in comparison to untreated cells was investigated by nano-liquid chromatography coupled with high-resolution tandem mass spectrometry. The schematic representation of the experimental design is reported in Fig. 1a. A high number of peptide groups (i.e. 14723) were used for the identification of 2134 proteins (Supplementary material, Table S1). The phosphoproteomic workflow enabled the identification of 430 phosphorylation sites from 510 phospho-peptides identified in at least one out of the two analyzed conditions (Supplementary material, Table S2A) mapping on 290 proteins (Supplementary material, Table S2B). Localization and best site probability for variable post-translational modifications within peptides were performed with the ptmRS tool of the Proteome Discoverer software that calculates individual probability values for each putatively modified site based on MS/MS data. Analysis of the phosphorylated amino acid distribution showed that 88% of phosphorylated sites were on serine, whereas 11% and 1% were threonine and tyrosine phosphorylation sites, respectively (Fig. 1b). These results are in line with previous literature data reporting that...
in human cells the overall levels of protein phosphotyrosines and phosphothreonines are lower than those of phosphoserines23,30,31.

In order to unravel the biological relevance of phosphorylation events triggered by W peptide stimulation, we focused our attention on a small number of peptides (53 mapping on 40 proteins) that were phosphorylated only in WKYMVm-stimulated cells (Table 1).

Functional analysis of identified phosphoproteins. The molecular function enrichment analysis of these proteins, performed through the Network analysis software, revealed that kinase binding, protein kinase binding, enzyme binding, proteins involved in transcription processes were among the most significantly enriched categories (Fig. 2a). We used PANTHER classification system to categorize FPR2-dependent phosphorylated proteins according to their known or postulated functions in biological processes. The analysis revealed that the phosphoproteins can be classified into eight groups and that Metabolic process represents the largest group in all the identified phosphorylated proteins (Fig. 2b). About 33% of the proteins of this group are involved in biosynthetic processes and we identified, among others, proteins involved in nucleotides, aminophosphonate, glyceralipid and cholesterol metabolisms, such as PRPP synthase-associated protein 2, choline-phosphate cytidylyltransferase A and hydroxymethylglutaryl-CoA synthase. The remaining 67% of proteins are involved in cellular metabolic processes, such as primary metabolism, organic substance metabolism and nitrogen compound metabolism. Biological regulation represents the second largest group in all the identified phosphorylated proteins (Fig. 2b). In particular, here we identified proteins involved in the regulation of cell cycle, cell division, apoptosis and transmembrane transport (Sororin, Promyelocytic leukemia protein PML, ATP-binding cassette). PANTHER analysis of the protein classes of the uniquely identified phosphoproteins showed that the nucleic acid binding proteins represent the largest group in all the phosphorylated proteins (Fig. 2c). About 77% of phosphoproteins of this group are RNA-binding proteins and about 23% are DNA-binding proteins. They play crucial roles in transcription, replication, recombination, DNA repair, and other nuclear functions. Notably, in this group we identified transcription and splicing regulatory proteins (RNA-binding motif protein, hRNP A3, hRNP C1 hRNP H1), proteins involved in the synthesis and maturation of ribosomes and translation initiation factors (Nucleolin, eIF-4B, eIF-4γ-1), cell cycle and DNA replication regulatory proteins (Rb, MCM2), histonic and non-histonic DNA-binding proteins (Histone H1.5, HMGI-C).
| Accessions | Description                                      | Peptide Sequence | Positions | ptmRS Probabilities |
|-----------|------------------------------------------------|-----------------|----------|---------------------|
| O06256-1 | PRPP synthase-associated protein 2             | [R].LGIAVHGIEAQAEDISLVDGHRSPMK. | [205–232] | S23: 100            |
| O95425–3 | Isoform SV3 of Supervillin                     | [R].SPSFQPPLSPAPR. [CV] | [261–276] | S10: 100            |
| O95747   | Serine/threonine-protein kinase OSRI           | [R].LHTKEDPGWGVEWDDEDEESEEGK. | [328–351] | S12: 100            |
| P04792   | Heat shock protein beta-1                      | [R].QLSQMSVEIR.  | [80–89]  | S3: 100             |
| P06400   | Retinoblastoma-associated protein              | [K].DREGPTDHLSACPLNLQNNHHTAAMDYLSFYRSPK. | [578–614] | S31: 98.09; S35: 100 |
| P06748   | Nucleophosmin                                   | [K].MQASIEKGSLPKVEAK. | [251–267] | S4: 100             |
| P07910-1 | Heterogeneous nuclear ribonucleoproteins C1    | [K].MESEGADDAAEEDLLDDDIDRDGDDELQ[IL]. | [238–271] | S10: 100            |
| P16401   | Histone H1.5                                    | [K].KATKSPAPK.   | [185–194] | S5: 100             |
| P23588   | Eukaryotic translation initiation factor 4B    | [R].ERHPSWRSETQER. | [402–412] | S5: 99.37           |
| P29590   | Protein PML                                     | [R].SPVYGSEVPLNNSVHГЧАЧЕПА, [V] | [530–555] | S1: 100             |
| P29966   | Myristoylated alanine-rich C-kinase substrate  | [K].SFKLGGSFK.   | [163–173] | S8: 100             |
| P31943   | Heterogeneous nuclear ribonucleoprotein H1     | [K].HTGPSNDTDANDGFR. | [99–114]  | S6: 100             |
| P38159-1 | RNA-binding motif protein, X chromosome         | [R].DVYLSRDPDDQYGTKDSSYR. | [204–223] | S5: 100             |
| P42167   | Lamina-associated polypeptide 2                | [R].AKTPVTLK.    | [206–213] | T3: 100             |
| P49006   | MARCKS-related protein                          | [K].KFSFKKPK.    | [91–99]   | S3: 100             |
| P49585   | Choline-phosphate cytidylyltransferase A       | [R].HKAAYDISEDEED. | [535–367] | S9: 100             |
| P51991-1 | Heterogeneous nuclear ribonucleoprotein A3     | [K].SSGSPYGGSOSGGSQGYGSRES. | [355–378] | S4: 100             |
| Q01518-1 | Adenyl cyclase-associated protein 1            | [R].SGKPFSAPKPQTPSPSKR. | [295–313] | T13: 99.99; S14: 50; S16: 50 |
| Q01581   | Hydroxymethylglutaryl-CoA synthase, cytoplasmic| [R].RPFTPDIDTDLDEGVGLVHISEATEHSPSK. | [469–498] | S27: 100            |
| Q04637-8 | Isoform 8 of eIF-4-4 gamma 1                   | [R].EALPPVSPLKAELSELEK. | [1225–1246] | S8: 100             |
| Q13442   | 28kDa heat- and acid-stable phosphoprotein     | [K].SLSDSEDEDEIDQYQQKR. | [57–75]  | S4: 100; S7: 100.   |
| Q13424-3 | Isoform 3 of Scaffold attachment factor B1     | [R].SYVSDKDYKEPR. | [601–612] | S4: 100             |
| Q8IU81   | Interferon regulatory factor 2-binding protein 1| [R].AGGASPAASSTAOQPTQHR. | [449–467] | S5: 100             |
| Q8NC51-1 | PAI1 RNA-binding protein 1                     | [R].GGGSHVWTGTVDELPTEYQKYQK. | [217–240] | S18: 96.8           |
| Q8WUD4   | Coiled-coil domain-containing protein 12       | [R].LKGQEDSLASAVIDAEQKTCDSQ. | [143–166] | S23: 100            |
| Q8WW12   | PEST proteolytic signal-containing nuclear protein | [K].TLSVAANFEDSEDEPEEMPEAKMR. | [106–131] | S14: 100            |
| Q8YJ36   | HMG box transcription factor BBX                | [R].TADGRVIPAGLTDKPEKQLQR. | [838–860] | S7: 99.99           |
| Q92538   | BFA-resistant GEF 1                           | [R].GTYTDSVEYTDHGEPKG. | [1315–1331] | S4: 99.46

Q96A00   | Protein phosphatase 1 regulatory subunit 14A   | [R].VLSKLQSPSR. | [10–19]   | S7: 99.72           |
| Q9HC35   | Echinoderm microtubule-associated protein-like 4| [R].ASPSQPQPQQPQLQR. | [143–158] | S4: 99.36           |
| Q9UDY2-7 | Isoform 7 of Tight junction protein ZO-2       | [R].SEEVPQHEESIKPSPEIR. | [983–1001] | S15: 99.98          |
| Q96FF9   | Sororin                                        | [R].SGFRAPEPTKRLR. | [15–28]   | S7: 99.99           |
| Q8NEF7-1 | ATP-binding cassette sub-family F member 1     | [K].KAEGQSEEEGEGESEELEESEGKADDHYALSK. | [223–255] | S6: 100             |
| Q9GZR7   | ATP-dependent RNA helicase DDX24               | [R].AQAVSEREEEEEGKSSPK. | [78–96]  | S5: 100; Th: 100; S9: 100 |
| Q9H6F5   | Coiled-coil domain-containing protein 86       | [R].APGPGRQHPKSPKPPKEPGTYGSGFGAK. | [185–212] | S4: 100             |

Continued
Phosphoproteins only identified in W peptide-stimulated CaLu-6 cells. Peptide sequence and positions within proteins together with accession numbers and descriptions are reported. Localization and best site probability for phosphorylation within peptides performed with the ptmRS tool of the Proteome Discoverer are also reported.

Table 1. Phosphopeptides only identified in W peptide-stimulated CaLu-6 cells. Peptide sequence and positions within proteins together with accession numbers and descriptions are reported. Localization and best site probability for phosphorylation within peptides performed with the ptmRS tool of the Proteome Discoverer are also reported.

To shed light on biological relationships among identified phosphoproteins, an interaction network map was constructed in silico by using the FunRich software (Fig. 3). To this aim, the dataset of all phospho-proteins identified by MS analysis (Fig. 3, grey nodes) was integrated with selected accession numbers of proteins known to be involved in response to FPR2 stimulation by W peptide (Fig. 3, red nodes). In particular, to investigate potential connection within the network and to point out the mutual interactions of identified proteins, the following nodes were included within the experimental dataset: Erk2 (P28482), Erk1 (P27361), PKCα (P17252), PKCβ (Q65565), p47phox (P14598), EGFR (P00533), PI3K (Q8WYR1), Akt (P31749), STAT3 (P40763)15,17,19. As shown, a large subset of proteins was mapped on a cluster converging on β-arrestin-1 (ARRB1), which plays a crucial role in GPCR signaling. This cluster also includes and displays connections with several phospho-proteins uniquely identified in WKYMVm-treated cells (Fig. 3, green nodes) such as HNRPNA3, TJP2, SVIL, TMPO, RB1.

Moreover, additional phosphoproteins were mapped on the network on core molecules known to be involved in FPR2 stimulation by W peptide, such as PRKCA and EGFR. HSP-27, MCM2, OSR1, Rb and MARCKS are selectively phosphorylated upon FPR2 stimulation by WKYMVm. FPR2 stimulation by WKYMvm exacerbates cell proliferation and the malignant phenotype of CaLu-6 cells, and its inhibition by PTX or WRWWWW significantly prevents these biological responses16. Therefore, we selected a subset of identified phospho-proteins (i.e. HSP-27, MCM2, OSR1, Rb and MARCKS) for further validation by western blot analysis, on the basis of their involvement in the control of cell proliferation and in signal transduction cascades triggered by FPR2 activation that we identified by MS/MS analysis.

HSP-27 (P04792) belongs to heat shock protein family and is expressed at basal level in many cell types and tissues. It is a chaperone32 with anti-inflammatory and immunomodulatory actions, including the release of the anti-inflammatory cytokine IL-1031. The protein is phosphorylated on three residues (Ser15, Ser78, and Ser82) by MAPKAP kinase 2/3/5 via the activation of p38MAPK pathway15. Other kinases, such as PKB, PKC, PKD, and PKG also independently phosphorylate HSP-2735. In fact, in SH-SY5Y human neuroblastoma cells, carbachol-dependent HSP-27 phosphorylation on Ser82 and Ser78 residues are prevented by PKC and p38MAPK inhibition, and phorbol esters-dependent HSP-27(Ser82) phosphorylation is reversed by PKC or PKD inhibitors46. Rac is also involved in HSP-27 phosphorylation, as demonstrated by the observation that the inhibition of Rac-guanine nucleotide exchange factor interaction reduces the collagen-induced HSP-27 phosphorylation on Ser15, Ser78 and Ser82 residues47. HSP-27 phosphorylation promotes the dissociation of HSP-27 multimers into monomers and dimers that can interact with other proteins. HSP-27 phosphorylation is highly dynamics and regulates several cellular processes in a context-specific manner. Indeed, phospho-HSP-27 is known to: (i) down-regulate the expression of intracellular glutathione and, in turn, the resistance against oxidative stress48; (ii) prevent Fas-induced apoptosis49; (iii) promote actin filament polymerization50 and (iv) modulate estrogen signaling51. In addition, a phospho-proteome analysis of tumors undergoing ischemia, HSP-27 results hyper-phosphorylated at the Ser82 residue suggesting the relevance of this site in ischemia42. Furthermore, in atherosclerotic lesions of hyper-cholesterolaemic rabbits a significant increase of HSP-27(Ser82) phosphorylation and pospho-p38MAPK was observed43. Consistent with previous reports reporting the induction of p38MAPK activity by WKYMVm13,44, our MS/MS analysis revealed FPR2-dependent phosphorylation of HSP-27 on Ser82 residue (Table 1). Accordingly, western blot experiments performed with an anti-phosphoHSP-27(Ser82) antibody showed an increased phosphorylation of HSP-27 in WKYMVm-stimulated cells, which is prevented by cells treatments with the FPR2 antagonist WRW4 or with PTX, which catalyzes the ADP-ribosylation of the αi subunits of the heterotrimeric G proteins. (Fig. 4a).

Stimulation of human monocytic U937 cells, or primary monocytes, or FPR2-transfected HEK293 cells with ANXA1 induces downstream phosphorylation of p38MAPK, MAPKAP kinase, and HSP-27. Pro-inflammatory
FPR2 agonists, such as LL-37 and Serum-Amyloid Alpha (SAA) are unable to phosphorylate HSP-27, suggesting that the activation of p38MAPK/MAPKAPK/HSP-27 pathway is a response triggered by pro-resolving, and not by pro-inflammatory ligands, upon FPR2 activation. Interestingly, FPR2 stimulation of monocytes with ANXA1 induces the release of IL-10, which is prevented by pretreatment with a p38MAPK inhibitor. Other members of the FPR family induce HSP-27 activation. In the human colonic epithelial cell line Caco2, physiological concentrations of N-fMLP significantly increase HSP-27 expression which is prevented by the FPR1 receptor antagonist BOC-FLFLF. In these cells N-fMLP activates p38MAPK and ERK1/2, but only p38MAPK inhibition prevents N-fMLP-dependent induction of HSP-27.

MCM2 (P49736) belongs to the protein family of “ATPases associated with diverse cellular activities” (AAA+). It is a component of the MCM2-7 complex, which is the putative replicative helicase necessary for DNA

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**Figure 2.** Bioinformatic analysis of WKYMVm-induced phosphoproteins. (a) Bar chart of the significantly enriched molecular functions of phosphoproteins uniquely identified in CaLu-6 cells following WKYMVm stimulation. The number of proteins belonging to the enriched gene ontology term and the p-values are reported on the graph. (b) PANTHER classification system of FPR2-dependent phosphorylated proteins in Biological Processes. (C) PANTHER analysis of the Protein Classes of the uniquely identified phosphoproteins.
replication initiation and elongation in eukaryotic cells. MCM2 can be phosphorylated by different kinases and these phosphorylation events are involved in DNA replication, cell cycle progression and checkpoints regulation. Cdc7/Dbf4 phosphorylates MCM2 and other MCMs in vivo and in vitro, on Ser27, Ser41, and Ser139 residues, whereas Cdk phosphorylates MCM2 but not on Ser139 residues. In human cells, Cdc7 is activated by its regulatory subunits Dbf4 and Drf1 and Cdc7/Dbf4 complex is directly involved in the initiation of DNA replication by targeting MCM2. Casein kinase 2 (CK2) and salt-inducible kinase 1 (SIK1) also phosphorylate MCM2 in vitro, but there is not evidence that CK2 is responsible for this phosphorylation in vivo. EGFR-and ERKs-dependent activation of CK2 phosphorylates phosphoglycerate kinase 1 (PGK1), resulting in PGK1/Cdc7 interaction. Cdc7-bound PGK1 converts the ADP in ATP thus removing ADP inhibition on Cdc7 and promoting MCM2 phosphorylation on Ser139. Previously, we demonstrated that WKYMVm stimulation of CaLu-6 cells induces EGFR transactivation and ERKs phosphorylation, and accordingly to EGFR-dependent activation of CK2/PKG1/Cdc7 cascade, our results potentially explain the observed FPR2-dependent phosphorylation of MCM2 on the Ser139 residue (Table 1). Western blot experiments performed with an anti-pMCM2(Ser139) antibody showed an increased phosphorylation levels of MCM2(Ser139) in FPR2-stimulated cells, and preincubation of CaLu-6 cells with WRW4 or PTX before W peptide stimulation prevent this phosphorylation (Fig. 4b).

The regulatory role of MCM2 in lung cancer has been extensively investigated in a integrate analysis of phospho-proteome and proteome of overexpressed and silenced MCM2 lung cancer cells. Such analysis demonstrated a phosphoMCM2-regulated functional network, suggesting that the deregulation of MCM2 phosphorylation is involved in lung cancer cell proliferation, cell cycle, and migration and that potential target cancer-specific phospho-proteins can be identified by the analysis of molecular interactions of phosphorylated MCM2. The role of phosphorylated MCM2 in cancer is also corroborated by a phospho-proteomic analysis of liver cell lines with different proliferation potential. The results show that MCM2 is hyper-phosphorylated in liver cancer in particular on a novel Thr27 phosphosite, but also on Ser139 residue. In these cells, MCM2 promotes cell proliferation via the regulation of high mobility group protein HMG-I/HMG-Y (HMG1) phosphorylation.
The oxidative stress–responsive kinase 1 OSR1 (O95747) is a serine/threonine-protein kinase involved in the regulation of the solute carrier 12 family of cation-chloride cotransporters and thereby in the modulation of cellular ion homeostasis, blood pressure, hearing, and kidney functions. OSR1 is activated by “with no lysine” (WNK) protein kinase family, which phosphorylates a Thr185 residue in the T-loop kinase domain, and Ser325 and Ser339 residues in the S-domain of OSR1. The role of OSR1(Ser325) and OSR1(Ser339) phosphorylations is unclear. Some evidence suggests that since the S-domain of OSR1 contains a WEW motif (aminoacids 336–338), essential for binding to the scaffolding protein MO25, the phosphorylation on serine residues adjacent to WEW motif (Ser339) could enhance binding to MO25. PI3K-Akt signaling activates the WNK-OSR1 cascade by Akt-dependent phosphorylation of WNK on Thr60, which is prevented by PI3K inhibitors. WNK3 is a direct target of Akt and is subjected to phosphorylation triggered by EGF-dependent PI3K-Akt pathway. Akt activity is regulated not only by PI3K phosphorylation in the activation loop (Thr308) but also by mammalian target of rapamycin complex 2 (mTORC2) phosphorylation in C-terminal hydrophobic motif (Ser473). mTORC2 also phosphorylates OSR1 on Ser339 residue, increasing OSR1 activity, and inhibition of mTORC2 does not prevent WNK activity, indicating that mTORC2 regulates OSR1 independently by WNK. Accordingly, OSR1 phosphorylation on Ser339 residue observed in WKYMVm-stimulated CaLu-6 cells suggests a link between FPR2-PI3K-Akt and Akt-mTORC2-OSR1 cascades. Consequently, western blot analysis performed with an anti-phospho antibody shows an increase of OSR1(Ser339) phosphorylation in FPR2-stimulated cells, which is prevented by WRW4 or PTX.

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The retinoblastoma tumor suppressor protein, Rb (P06400), negatively regulates entry into S phase. The binding of Rb to E2F prevents the release of E2F, thereby inhibiting the entry of the cell cycle. Therefore, Rb acts as a cell cycle inhibitor and is activated by multisite Cdk-dependent phosphorylation in response to positive growth signals. The Rb protein consists of several domains (Rhb, RbD1, Rbc), as well as a large loop within the pocket domain (RbPL). Cdk-dependent Rb phosphorylation occurs at 13 consensus sites which include all these regions. Unphosphorylated Rb on Ser608 residue has growth suppressive activity and specifically contact the COOH-terminus of MCM4. Phosphorylation of Ser608 in RbPL is sufficient for partial inhibition of E2F.

Figure 4. Validation of FPR2-dependent phosphoproteins. Western blot and densitometric analyses of at least three independent experiments. Fifty micrograms of whole lysates were purified from control cells and from 24 hours serum-starved CaLu-6 cells stimulated with 10 μM WKYMVm for 5 minutes, in the presence or absence of 10 μM WRW4 (WRW4) or 100 ng/mL PTX. Lysates were resolved on 10% SDS-PAGE and hybridized with anti phospho-selective antibodies specific for p-HSP-27(Ser82) (A); p-MCM2(Ser139) (B); p-OSR1(Ser339) (C); p-Rb(Ser608) (D) and p-MARCKS(Ser170) (E). An anti-tubulin antibody was used as loading proteins control.
binding, whereas Ser608/Thr373 phosphorylation produces unique conformational changes of the Rb structure that result in the allosteric and complete inhibition of the E2F binding. Ser608/Ser612 phosphorylation induces an intramolecular association between RbPL and the pocket domain. This association obstructs the E2FTD binding site in the pocket, preventing the simultaneous binding of phosphorylated RbPL and E2FTD.

Rb is not phosphorylated on Ser608 residue and is tethered to nuclear structures in CD34+ hematopoietic progenitor cells, suggesting that it has growth-suppressive activity, whereas Rb(Ser608) is phosphorylated by CDK4/6 complexes in acute lymphoblastic leukemia cells. The role of Rb(Ser608) phosphorylation in the control of the progression of the cell cycle is further confirmed in a subset of tumor-derived cell lines, where conversion of unphosphorylated Rb in a phosphorylated form does not arise. In these defective cells, a lack of dephosphorylation of Rb(Ser608) is observed. Several members of PP1 phosphatase family are implicated in the dephosphorylation, but there is no evidence for defective or deregulated expression of catalytic PP1 cores. We decided to validate the identified Rb(Ser608) phosphorylation on the basis of its role in the control of cell cycle and, in western blot analysis, we observed that an anti-pRb(Ser608) antibody confirmed this phosphorylation in FPR2-stimulated cells, which was prevented by preincubation with WRW4 or PTX (Fig. 4d).

MARCKS (P29966) is a membrane-binding protein and is the main substrate of PKC in many cell types. It binds calmodulin (CaM), actin, synapsin and is involved in cell migration, regulation of cell cycle, actin-filament reorganization, adhesion, cytoskeletal control and neurosecretion. PKC-dependent phosphorylation on Ser159, Ser163, and Ser170 residues within PSD increases MARCKS detachment from the membrane and removes PIP2 association. PKC-dependent phosphorylation of MARCKS reduces its affinity for CaM, suggesting that MARCKS could represent the storage of CaM and that phospho-MARCKS could play a crucial role in regulating CaM availability within the cell. MARCKS is preferentially phosphorylated by PKCo but also by PKCd and PKCe, which phosphorylate Ser159, Ser163, and Ser170 residues in the PSD of human MARCKS. These phosphorylations mask positive charges on the protein which, in turn, segregates from the plasma membrane to the cytoplasm. Rho kinase (ROCK) also phosphorylates PSD of MARCKS and, in mouse bone granulocytes, W
peptide stimulation triggers ROCK activation\(^7^5\). We previously observed the ability of FPR2 to activate PKC\(\alpha\) and PKC\(\delta\) in several cell types\(^3^,1^7,1^9\) and, accordingly, our phospho-proteomic analysis identified the FPR2-dependent MARCKS(Ser170) phosphorylation (Table 1). In protein extracts from WKYMVm-stimulated cells, the selective phosphorylation of MARCKS(Ser170) was detected by a specific antibody and this phosphorylation is prevented by the FPR2 antagonist and by Gi protein inhibition (Fig. 4e). These results are in line with previous evidence reporting that stimulation of FPR1 with its agonist N-fMLP triggers p47 phox and MARCKS phosphorylation\(^7^6\).

In the lung, phospho-MARCKS plays a key role in the control of mucin secretion and inflammation. Higher phospho-MARCKS levels are correlated with shorter overall survival of lung cancer patients, suggesting a key role for MARCKS in lung cancer and a therapeutic strategy for inhibiting PKC activity\(^7^7\).

We also used specific inhibitors to determine the interconnection between some of the signalling readouts identified. PKC and MAPK represent two nodes in the interaction network analysis of proteins identified by LC-MS/MS (Fig. 3). Therefore, we preincubated cells with Rottlerin or PD098059 prior WKYMVm stimulation, and in western blot experiments we observed that the inhibition of PKC\(\delta\), which is activated upon FPR2 stimulation\(^3,1^7,1^9\), prevents WKYMVm-induced MARCKS(Ser170) phosphorylation (Fig. 4e). These results are in line with previous evidence reporting that stimulation of FPR1 with its agonist N-fMLP triggers p47 phox and MARCKS phosphorylation\(^7^6\).

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**FPR2 stimulation with anti-inflammatory agonists induces HSP-27, OSR1, Rb and MARCKS phosphorylation.** Several data support the development of WKYMVm as a novel and effective anti-inflammatory therapeutic agent. It shows pleiotropic, immunomodulatory, anti-apoptotic, and anti-inflammatory effects in several pathological conditions\(^8^\), inhibits the production of pro-inflammatory

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**Figure 6.** HSP-27, OSR1, Rb and MARCKS phosphorylation depends on ANXA1 stimulation. Serum-starved CaLu-6 cells were stimulated or not with 10 nM ANXA1 for 5 minutes, in the presence or absence of 10 \(\mu\)M WRW4 or 100 ng/mL PTX. Fifty micrograms of whole lysates were purified, resolved on 10% SDS-PAGE and hybridized with anti-p-HSP-27(Ser82) (a); anti-p-OSR1(Ser339) (b); anti-p-Rb(Ser608) (c); anti-p-MARCKS(Ser170) (d), and anti-p-MCM2(Ser139) antibodies. Proteins were detected by autoradiography and bands densitometry was estimated using a Discover Pharmacia scanner. The filters were reprobed with an anti-tubulin antibody to normalize the amount of loaded proteins. Data are representative of three independent experiments.
cytokines (TNF-α, IL-1β, and IL-6) and enhances the levels of anti-inflammatory cytokines (IL-10 and TGFβ)⁷⁹. FPR2 can transmit both pro- and anti-inflammatory signals, and this duality was originally established by the nature of the ligands. In general, short and flexible structures, like that of formylated peptides, activate a pro-inflammatory cell response, while ANXA1 and LXA4 are well-known anti-inflammatory FPR2 ligands. The shift between pro- and anti-inflammatory cell responses is caused by conformational modifications of FPR2 upon ligand binding, but are not merely due to ligand-receptor interaction. FPR2 can dimerize and this conformational change could account for its biological functions. Anti-inflammatory agonists enhance the formation of FPR2 homodimers and the release of inflammation-resolving cytokines, while inflammatory ligands, such as SAA, do not cause receptor homodimerization. Heterodimers of FPR2 with other members of the FPR family can transduce proapoptotic signals⁴³. We stimulated CaLu-6 cells with two anti-inflammatory agonists and, in western blot experiments, we observed that ANXA1 and LXA4 induce HSP-27(Ser82), OSR1(Ser339), Rb(Ser608), and MARCKS(Ser170) phosphorylation, which was prevented by PTX and WRW4 (Figs. 6A–D and 7A–D). However, no significant variation was detected in MCM2(Ser139) phosphorylation (Figs. 6E and 7E), suggesting that the latter only depend on WKYMVm-stimulation. ANXA1 stimulation of FPR2-expressing cells induces the phosphorylation of p38MAPK/MAPKAP/HSP-27 cascade⁴⁵, but in these cells, the phosphorylation of HSP-27 on a Ser82 residue has not been demonstrated. Anti-inflammatory ligands increase the formation of FPR2 homodimers and/or FPR1/FPR2 heterodimers and, in turn, the release of anti-inflammatory cytokines⁴⁵. Since CaLu-6 cells don't express FPR1/FPR2, but not FPR2/FPR3 heterodimerization.

**Conclusions**

FPR2 was initially detected in phagocytic leukocytes, but the receptor was subsequently identified in other cell types and tissues, where it is involved in many physio-pathological processes and where it shows further functions than those normally exerted in polymorphonuclear cells. For instance, FPR2 promotes the malignancy of colon cancer⁸¹, triggers c-Met and EGFR transactivation in PNT1A and CaLu-6 cells⁶¹⁹, respectively, and

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**Figure 7.** LXA4 induces HSP-27, OSR1, Rb and MARCKS phosphorylation. CaLu-6 cells were serum-starved for 24 hours and stimulated or not with 1 μM LXA4 for 5 minutes. Cells were also preincubated with 10 μM WRW4 or 100 ng/mL PTX before LXA4 stimulation. Whole lysates (50 μg) were resolved on 10% SDS-PAGE and hybridized with anti-p-HSP-27(Ser82) (a); anti-p-OSR1(Ser339) (b); anti-p-Rb(Ser608) (c); anti-p-MARCKS(Ser170) (d), and anti-p-MCM2(Ser139) antibodies. Proteins were detected by autoradiography and bands were estimated from densitometric scanning. An anti-tubulin antibody was used to normalize the amount of loaded proteins. Results are representative of three independent experiments performed in triplicate.
elicits NOX2-dependent superoxide generation in IMR90 human fibroblasts, suggesting a specific role of FPR2 in non-phagocytic cells. CaLu-6 is a human lung anaplastic cancer cell line which expresses high levels of EGFR as well as a biological functional FPR2. We already demonstrated that stimulation with W peptide exacerbates the malignant phenotype of these cells through a molecular mechanism which involves EGFR tyrosine trans-phosphorylation, p47phox phosphorylation, NADPH-oxidase–dependent superoxide generation, and c-Src kinase and STAT3 activity. FPR2 inhibition significantly prevents cell proliferation and the malignant phenotype, suggesting the potential role of FPR2 signalling and, in turn, of phosphorylated proteins as drug targets in lung cancer epithelial cells.

The investigation of the complex and dynamic regulation systems of protein phosphorylation represents an attractive field of proteomic research. The outline of the cellular phosphorylation status is crucial for the identification of intracellular signalling pathways and, in turn, for the understanding of the molecular mechanisms responsible for cancer and other human disorders. Here we report the first phospho-proteomic analysis of FPR2-stimulated CaLu-6 cells that allowed the overall identification of 290 phosphorylated proteins; among them, we found 53 unique phosphopeptides that mapped on 40 proteins with crucial roles in metabolic processes and biological regulation. In particular, we identified FPR2-induced phosphorylated proteins involved in the control of cell cycle progression, cell division and apoptosis, and thereby implicated in molecular mechanisms of cellular invasion and metastasis. Interestingly, most of these phospho-proteins are transcriptional factors or are regulators of replication, recombination, transcription, splicing, translation processes and DNA repair. The heterogeneity and complexity of phospho-protein patterns are indicative of the activity of multiple signal transduction pathways triggered by FPR2 stimulation. Several novel phosphosites have been identified in this study, which represent an important basis for further functional studies that may contribute to the identification of new signalling networks. However we cannot exclude that the phosphorylation pattern that we observed is a general phenomenon that occurs during FPR2 activation in all FPR2-expressing cells, rather than specific of lung epithelial carcinoma CaLu-6 cells.

FPR2 can transmit both pro- and anti-inflammatory signals and this ability is determined by the nature of the ligands and by the formation of higher-order structures. For instance, FPR2 can homodimerize or to heterodimerize with FPR1 or FPR3. Pro-resolving ligands, such as ANXA1 and Ac2-26, increase the formation of FPR2 homodimers and/or FPR1/FPR2 heterodimers, which led to the release of inflammation-resolving cytokines, while pro-inflammatory agonists do not cause homodimerization. The receptor oligomerization is not limited to the FPR family members but also includes the scavenger receptor MARCO and is connected to agonist-evoked modifications in ERKs phosphorylation and cAMP levels. We can exclude FPR1/FPR2 heterodimerization in CaLu-6 cells since these cells don’t express FPR1, but we cannot rule out the FPR2/FPR3 heterodimerization. Our results strongly suggest a potential FPR2 homodimerization upon WKYMVm stimulation, since FPR2 oligomerizes with FPR1 or FPR3. Pro-resolving ligands, such as ANXA1, which increases the formation of FPR2 homodimers, induces the phosphorylation of HSP-27 and of other validated proteins (Fig. 6).

The identification of the intracellular signalling pathways triggered by FPR2, as well as of downstream phosphorylated proteins, will provide a better understanding of the role of FPR2 in non-phagocytic cells in physiological and pathological conditions. Furthermore, these results could provide new insights into FPR2 signalling as possible drug targets in human diseases, including inflammatory disorders, infections, and cancer, that involve this receptor.

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F.C., P.V.P. and R.A. designed the experiments. F.C., R.R., M.C., A.C. and C.Z. performed the experiments. A.C., G.E., P.V.P. and R.A. analysed the data and wrote the manuscript. A.C., P.V.P. and R.A. obtained funding

**Competing interests**

The authors declare no competing interests.

**Additional information**

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