Evaluation of low doses BPA-induced perturbation of
glycemia by toxicogenomics points to a primary role of
pancreatic islets and to the mechanism of toxicity

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Epidemiologic and experimental studies have associated changes of blood glucose homeostasis to Bisphenol A (BPA) exposure. We took a toxicogenomic approach to investigate the mechanisms of low-dose (1 × 10⁻⁹ M) BPA toxicity in ex vivo cultures of primary murine pancreatic islets and hepatocytes. Twenty-nine inhibited genes were identified in islets and none in exposed hepatocytes. Although their expression was slightly altered, their impaired cellular level, as a whole, resulted in specific phenotypic changes. Damage of mitochondrial function and metabolism, as predicted by bioinformatics analyses, was observed: BPA exposure led to a time-dependent decrease in mitochondrial membrane potential, to an increase of ROS cellular levels and, finally, to an induction of apoptosis, attributable to the bigger Bax/Bcl-2 ratio owing to activation of NF-κB pathway. Our data suggest a multifactorial mechanism for BPA toxicity in pancreatic islets with emphasis to mitochondria dysfunction and NF-κB activation. Finally, we assessed in vitro the viability of BPA-treated islets in stressing condition, as exposure to high glucose, evidencing a reduced ability of the exposed islets to respond to further damages. The result was confirmed in vivo evaluating the reduction of glycemia in hyperglycemic mice transplanted with control and BPA-treated pancreatic islets. The reported findings identify the pancreatic islet as the main target of BPA toxicity in impairing the glycemia. They suggest that the BPA exposure can weaken the response of the pancreatic islets to damages. The last observation could represent a broader concept whose consideration should lead to the development of experimental plans better reproducing the multiple exposure conditions.

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In the past decade, a huge effort has been made to identify xenobiotics that represent a risk factor for glucose homeostasis and diabetes development, impairing hepatocytes function, insulin production/response and, finally, reducing the β-cell mass.¹⁻³ Loss of β-cells is typically associated to type 1 diabetes (T1D), even though their failure has been involved in the pathogenesis of obesity-associated type 2 diabetes (T2D).⁴ Estrogens and xenoestrogens have a direct role in the regulation of glucose homeostasis, enhancing the secretion of insulin and affecting β-cells survival via non-genomic pathway, through estrogen receptor α (ERα) and β (ERβ) or G protein-coupled estrogen receptor (GPR30).⁵⁻⁸

Bisphenol A (BPA), a well-characterized xenoestrogen, is described as risk factor for development of T2D.⁸⁻¹² It is used in the manufacture of polycarbonates and epoxy resins. Its polymers, not toxic, are used in food contact materials. Monomers, exerting the estrogenic activity, can seep into the water or food following breakdown of polymers.¹³ The EFSA considers diet as the major source of BPA exposure.¹⁴ Exposure to BPA is nearly ubiquitous with concentration in human serum ranging from 0.2 to 1.6 ng/ml and it accumulates in fat.¹⁵⁻¹⁷ In the last decade, the interest for potential effects of BPA on metabolic disorders is on the rise. Indeed, a positive correlation between diabetes and BPA urinary levels was found combining the data from three epidemiological NHANES studies¹⁸ while other reports retrieved different degree of associations.² However, increasing in vitro and in vivo evidences suggest that BPA affects functions of pancreatic islets.¹⁹⁻²⁶ It has been shown that exposure to low doses of BPA, in vitro, could cause an increase of insulin content and release in the islets throughout ER pathways or CREB activation.¹⁹ Indeed, the in vivo exposure to 10 μg/kg/day or 100 μg/kg/day altered the glycemic and insulin curve, favouring insulin resistance.²⁵ In murine pancreatic islets, BPA has been shown to act similarly to E₂ binding membrane-bound ER and exerting non-genomic actions.²⁶ In the same

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Abbreviations: BPA, bisphenol A; CCCP, carbonyl cyanide 3-chlorophenylhydrazone; CREB, cAMP response element-binding protein; E2, 17β-estradiol; EDTA, ethylenediaminetetraacetic acid; EGTA, ethylene glycol tetraacetic acid; ER, estrogen receptor; FBS, fetal bovine serum; FOXO, forkhead domain factors; GPR30, G protein-coupled estrogen receptor; HBSS, Hank’s Balanced Salt Solution; IκB, NF-kappa-B inhibitor alpha; IPA, Ingenuity Pathway Analysis; NAC, N-acetylcysteine; NF-κB, nuclear factor kappa-light-chain-enhancer of activated B cells; NHANES, National Health and Nutrition Examination Survey; PBS, phosphate-buffered saline; ROI, region of interest; ROS, reacting oxygen species; STZ, streptozoticin; T1D, diabetes type 1; T2D, diabetes type 2

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context, exposure to low dose of BPA (1 × 10^{-9} M) induced the increase of intracellular [Ca^{2+}], insulin release and, finally, the transcription of CREB-dependent genes (e.g., insulin gene). Furthermore, Lin et al. demonstrated that low concentration of BPA could cause mitochondrial dysfunction and cell death in INS-1E, a rat insulinoma cell line.

The liver also has a relevant role in maintenance of the glucose balance through the regulation of its storage and release. Moreover, this organ is the major site of BPA metabolism and exposure to BPA has been associated with abnormalities of liver function and hepatic damage. In vivo and in vitro studies documented in the hepatocytes an altered gene expression after exposure to different doses of BPA. An increased apoptosis was also detected in hepatocytes after exposure to reference dose of 50 μg/kg/day, from gestation day 0 to the postnatal day 21 (ref. 32). Perinatal exposure to BPA has been linked to hypermethylation of glucokinase promoter and to inhibition of its gene expression that could contribute to the development of insulin resistance in the adulthood.

The summarized results suggest a clear role of BPA in the impairment of glucose homeostasis. They sketch the mechanisms of BPA toxicity even if the molecular aspects are not completely defined. The use of the ‘omics’ approaches has been strongly suggested to explore the mechanisms of toxicity exerted by BPA and other endocrine disrupting chemicals. Indeed, the gene expression profiles contain a significant amount of information on the current biological conditions leading to a better understanding of related phenotypic and molecular changes. Therefore, we chose the transcriptomic approach to underscore the key molecular events through which BPA impairs glucose homeostasis in primary cells. Primary ex vivo cell cultures have been preferred to immortalized cell lines to preserve different cell types and their interactions within the pancreatic islets and to in vivo studies to reduce the number of enrolled animals.

Here we report the results of a study aimed to investigate the mechanisms of toxicity of environmental dose of BPA (1 × 10^{-9} M) on cultured murine primary hepatocytes and pancreatic islets by a toxicogenomic approach. Our work do not identify any significant transcriptional effects in hepatocytes but confirms the pivotal role of mitochondrial dysfunction in the apoptosis induced by BPA exposure of pancreatic islets. We report the map of the slight transcriptome alterations whose summary results in a reduced viability of the pancreatic islets. Importantly, accordingly to our data, BPA exposure impairs the recovery of islet cells from damages, thus suggesting a mechanism for the diabetogen role of BPA.

Results

Toxicogenomics of murine primary hepatocytes and pancreatic islets exposed to low dose of BPA. To molecularly define the effects that BPA exerted on glucose homeostasis, we analyzed the transcriptional response in two tissues involved in glucose metabolism: liver and pancreas. Cultures of murine primary hepatocytes and pancreatic islets were prepared and exposed to low dose (1 × 10^{-9} M) BPA for 48 h. The time was fixed to analyze the effects on healthy cultured primary cells (see Supplementary Figure S1). Surprisingly, ex vivo exposure to 1 × 10^{-9} M BPA did not affect the transcriptome of hepatocytes (Figure 1a). Conversely, a small group of inhibited genes was identified in islets, as shown by the Volcano Plot (Figure 1b) and by the Heatmap (Figure 1c). Twenty-nine genes were differentially expressed. They are listed in Table 1, where the fold changes of microarray and qRT-PCR are reported. Furthermore, for six selected genes, we confirmed the specific inhibition for islets as it was not retrieved in the hepatocytes (Figure 1d). By Ingenuity Pathway Analysis (IPA), some of the inhibited genes were involved in two deregulated Canonical Pathways both pertaining to mitochondrial function: oxidative phosphorylation and mitochondrial dysfunction (Log B-H P-value, 2.12). Transcript level of some of these genes was determined to assess the timing of the inhibition. We selected genes encoding components of respiratory chain complexes (Uqcrb and Ndufs4), proteins involved in cell detoxification (Gpx3, Sod2 and Zfand2A), proteins involved in ATP production and/or in insulin exocytosis process (ATP1b1, ATP6v1f and Vapa). Isocitrate-tRNA synthetase (lars) was selected as defects in aminoacetyltransferase of tRNA were associated to the impairment of cytochrome c oxidase activity. To this aim, islets were exposed to 1 × 10^{-9} M BPA for 24 h and 48 h. The data confirmed the inhibition of selected transcripts at 48 h, starting already at 24 h for some of them (Figures 2a and d). The inhibition of ATP1b1, Vapa and Zfand2a was stronger at 24 h than 48 h (Figures 2b and d). We did not observe any significant modification in the protein levels of some of them (data not shown), maybe owing to the slight reduction of their transcripts with respect to their high cellular content.

Mitochondrial dysfunction and cell viability in pancreatic islets exposed to low dose of BPA. Starting from the bioinformatic prediction of altered mitochondrial function, we decided to assess the intracellular oxidative stress and mitochondrial activity in the treated islets to confirm functionally the microarray results. The exposure was performed on dispersed pancreatic islet cells, treated with low-dose (1 × 10^{-9} M) BPA for different times. Although we did not find a clear reduction in the protein levels of the deregulated transcripts, we could detect a clear and time-dependent increment of intracellular ROS level after 12 h exposure (fold change 1.79, Figure 3a) and a subsequent impairment of mitochondrial membrane potential at 18 h (0.76) and 24 h (0.74) of treatment (Figure 3b). The BPA effects on islet survival were analyzed by MTT assay. Higher doses (1 × 10^{-4} M and 1 × 10^{-6} M) were also tested. BPA affected the cellular viability of dispersed islets in a dose-dependent manner (Figure 3c). The highest BPA dose was already toxic after 24 h with an activity similar to 25 mM glucose (0.64 and 0.48 at 24 h and 48 h, respectively). Exposure to lower doses of BPA (1 × 10^{-6} M and 1 × 10^{-9} M) reduced the cell viability only at 48 h (0.70 and 0.76, respectively).

The above-reported data suggested that BPA exposure could damage islet cells throughout alteration of mitochondrial function and increasing ROS cellular level. Both aspects have been implied in the islet viability. We assessed the apoptosis at single cell level by TUNEL assay performed on
exposure to $1 \times 10^{-9}$ BPA was similar to the one exerted by menadione (23.42 and 23.42%, respectively). Noteworthy, the apoptotic effect at $1 \times 10^{-9}$ M BPA exposure was further confirmed assessing the number of apoptotic cells, in co-exposed cells, when compared with BPA-only-treated cells (1x10$^{-6}$ M BPA: Vehicle 41.8%, NAC 29.2%, BMS 32.7%; 9 M BPA: Vehicle 26.2%, NAC 18.7%, BMS 21.6%). As reported in Figure 4c, both inhibitors deeply affected the $Bax$ mRNA increase due to BPA. This molecular result was phenotypically confirmed assessing the ability of NAC and BMS-345541 to revert the BPA-dependent apoptosis in treated cells. To this aim, TUNEL assay was performed on dispersed islets co-exposed for 48 h to BPA (1x10$^{-6}$ M and 1x10$^{-9}$ M) and NAC or BMS-345541. As reported in Figure 4c, both inhibitors significantly reduced the number of apoptotic cells, in co-treated cells, when compared with BPA-only-treated cells (1x10$^{-4}$ M BPA: Vehicle 41.8%, NAC 29.2%, BMS 32.7%; 1x10$^{-9}$ M BPA: Vehicle 26.2%, NAC 18.7%, BMS 21.6%), pointing to a clear role of ROS induction and NF-kB activation in the process.

Cellular pathways involved in pancreatic islet damage following low-dose BPA exposure. Although BPA best-known targets are ERs, BPA is able to exert its activity through other signaling pathways, as NF-κB. The IPA analyses predicted several BPA targets as Forkhead Domain (FOXO, $P$-value 0.02) and NF-κB ($P$-value 0.04), among the others. We verified through a bioinformatics tool, Genomatix, if ER, NF-kB and FOXO binding sites were predicted in the $Bax$ promoter (Figure 4a) as well as in the promoters of the other previously selected genes. They were found in all the analyzed promoters (data not shown). Given the role of $Bax$ in the apoptosis, we focused on this gene and assessed the effects, on BPA activity, of inhibitors of the cited pathways: BMS-345541 (NF-κB inhibitor), LY-294002 (PI3K/AKT inhibitor, a regulator of FOXO) and ICI-182780 (ERα antagonist). The role of NAC (a ROS inhibitor) was also assessed. To this aim, purified islets were co-exposed for 48 h to $1 \times 10^{-9}$ M BPA and different inhibitors. The co-exposure of the islets to BPA and ICI-182780 or LY-294002 did not significantly affect the upregulation of $Bax$ (Figure 4b). Even though NAC and BMS-345541 exposure, slightly and not significantly, inhibited $Bax$ transcript, both inhibitors deeply affected the $Bax$ mRNA increase due to BPA. This molecular result was phenotypically confirmed assessing the ability of NAC and BMS-345541 to revert the BPA-dependent apoptosis in treated cells. To this aim, TUNEL assay was performed on dispersed islets co-exposed for 48 h to BPA (1x10$^{-6}$ M and 1x10$^{-9}$ M) and NAC or BMS-345541. As reported in Figure 4c, both inhibitors significantly reduced the number of apoptotic cells, in co-treated cells, when compared with BPA-only-treated cells (1x10$^{-4}$ M BPA: Vehicle 41.8%, NAC 29.2%, BMS 32.7%; 1x10$^{-9}$ M BPA: Vehicle 26.2%, NAC 18.7%, BMS 21.6%), pointing to a clear role of ROS induction and NF-kB activation in the process.

The activation of the NF-kB pathway by low dose BPA exposure was further confirmed assessing the cellular level of IkB-α that has to be degraded to allow NF-kB nuclear translocation. To this aim, whole cellular lysates were
Figure 2  Time-dependent effects of BPA on transcript inhibition. Level of transcripts was determined by qRT-PCR after exposure to $1 \times 10^{-9}$ M BPA for 24 h and 48 h. The selected genes were grouped by functional categories: respiratory chain subunits, Uqcrb and Ndufs4 (a); ATP-dependent pump subunits, ATP1b1 and ATP6v1f (b); ROS detoxification, Gpx3 and Sod2 (c); protein synthesis and degradation, Vapa, Iars, Zfand2a (d). Data are reported as fold change values. The results are expressed as the mean± SD of three independent experiments (N=3). *P-value <0.05; **P-value <0.01; ***P-value <0.001 compared with vehicle-only-treated islets.
prepared from pancreatic islets treated with BPA (1 × 10⁻⁹ M and 1 × 10⁻⁶ M) and IkB-α cellular content assayed by western blotting. The time points, 12 h, 18 h and 48 h, were chosen according to ROS level, mitochondrial potential and gene expression assays, respectively. As shown in Figure 4d, IkB-α protein level was decreased after 12 h of exposure at 1x10⁻⁶ M BPA, in correspondence with the peak of ROS production. At later time points, 18 h (Figure 4d) and 48 h (Figure 4e and Supplementary Figure S3), it increased, probably as a result of the NF-κB activation. Indeed, IkB-α is a well-known NF-κB target gene and the increase in its transcript is detectable by qRT-PCR already at 12 h of exposure (Figure 4f). At 1x10⁻⁶ M BPA exposure, we observed a degradation of IkB-α at a later time (18 h), maybe due to more stressing condition or an inverse dose-effect response already described for BPA. The NF-κB activity increase was further documented by assessing nuclear p65/Rel A in the nuclei of BPA-treated islet cells by whole-mount immunohistochemistry (Supplementary Figure S4).

Low dose exposed murine pancreatic islets show an impaired response to glucose stress in vitro and in vivo. The described results suggested that BPA toxicity on murine ex vivo cultured pancreatic islets was exerted altering the mitochondria functionality and lowering the cellular viability. As no massive effects were detected on cell viability, we speculated that BPA could act, mainly, altering the islet response/recovery from an injury, as glucose overload. To test this hypothesis, we assessed the ATP production in islet cells, dispersed and cultured, co-exposed to BPA (1 × 10⁻⁹ M) and glucose (25 mM). As reported in Table 2, the cells co-treated with glucose and 1 × 10⁻⁶ Mo r 1x10⁻⁹ M BPA showed a reduction of ATP content up to 0.42 and 0.35 fold change versus the glucose-only-treated islets. Furthermore, the treatment with both 1 × 10⁻⁶ M and 1 × 10⁻⁹ M BPA for 48 h reduced the secretion of insulin in response to 1 h stimulation with 16 mM glucose (fold change 0.65 and 0.73, respectively; Figure 5a). We confirmed the results testing the ability of the treated islets to restore the glycemia after their transplantation in diabetic mice. For in vivo experiment, primary islets were prepared and cultured for 48 h with 1 × 10⁻⁹ M BPA or vehicle alone. To ensure the restoration of the normal glycemia level despite the apoptosis induced by BPA treatment, 400 BPA-treated and control islets were transplanted in STZ injected mice. As shown in Figure 5b, the transplant with BPA pre-treated islets was unable to restore normal glycemic level neither in BPA treated (Gr.2) nor in normal water (Gr.3)-administered mice at any
Figure 4  Pathways involved in BPA apoptosis triggering. (a) Schematic representation of transcription binding sites for fork head domain factors (FKHD), nuclear factor kappa B (NF-κB) and estrogen responsive elements (ERE) predicted by Genomatix software in Bax gene promoter fragment −1000/+50 bp; (b) qRT-PCR analyses of Bax transcript in murine pancreatic islets cultured in the presence or absence of 1 × 10^{-9} M BPA, 3 mM NAC, 2 × 10^{-5} M BMS-345541, 1 × 10^{-5} M LY-294002 or 1 × 10^{-5} M ICI-182,780, for 48 h. Data are reported as fold change values. *P-value < 0.05; **P-value < 0.01; ***P-value < 0.001 compared with BPA-only-treated islets; (c) IF-TUNEL staining of cells co-exposed to BPA and ROS and NF-κB inhibitor. Dispersed islet cells were cultured in chamber slides in the presence or absence of 1 × 10^{-4} M BPA, 1 × 10^{-9} M BPA, 3 mM NAC, 2 × 10^{-5} M BMS-345541 for 48 h and then visualized under a fluorescence microscope. Data are reported as percentage of TUNEL-positive cells/total cell number (DAPI staining). *P-value < 0.05; **P-value < 0.01; ***P-value < 0.001 compared with BPA-only-treated islets; (d) IKK-α protein was determined in whole cellular extracts prepared from intact murine pancreatic islets cultured in the presence or absence of 1 × 10^{-4} M BPA and 1 × 10^{-9} M BPA for 12 h and 18 h by western blot; (e) protein quantification of IKK-α in murine pancreatic islets as already reported (d and Supplementary Figure S3). The signal intensity was determined with ImageJ software and normalized on β-actin protein signal intensity. Data are reported as fold change values. *P-value < 0.05; **P-value < 0.01; ***P-value < 0.001 compared with vehicle-only-treated islets; (f) qRT-PCR quantification of IKK-α transcript in murine pancreatic islets. Level of transcripts was determined by qRT-PCR after exposure to 1 × 10^{-9} M BPA for 12 h. Data are reported as fold change values. *P-value < 0.05; **P-value < 0.01; ***P-value < 0.001 compared with vehicle-only-treated islets. All the results are expressed as the mean ± SD of three independent experiments (N = 3)

Table 2  BPA affects ATP content in pancreatic islets cultured in the presence of 25 mM glucose

| BPA      | Glucose 25 mmol/l | Normalized on vehicle-treated cells | Normalized on glucose-treated cells |
|----------|-------------------|------------------------------------|-------------------------------------|
|          |                   | Relative luminescence level fold change | St. Dev. | P-value | Relative luminescence level fold change | St. Dev. | P-value |
| Vehicle  | −                  | 1.000                             | 0.247  |         | 1.000                             | 0.474  |         |
| 10^{-6} M| −                  | 0.983                             | 0.330  | 0.922   | 0.418                             | 0.177  | 0.018*  |
| 10^{-9} M| −                  | 1.047                             | 0.244  | 0.745   | 0.348                             | 0.137  | 0.009** |
| −        | +                  | 0.261                             | 0.314  | 0.001   | 1.000                             | 0.474  |         |
| 10^{-6} M| +                  | 0.088                             | 0.107  | 0.0001  | 0.102                             | 0.103  | 0.0001  |
| 10^{-9} M| +                  | 0.102                             | 0.103  |         | 0.418                             | 0.177  | 0.018*  |

Data are reported as fold change values calculated as ratio between average luminescence/total protein content in treated and control cells. *P-value < 0.05; **P-value < 0.01; compared with glucose-only-treated islets
time. We also found a loss of weight in the BPA-treated animals significantly greater in treated mice than controls (data not shown). As already shown, the BPA exposure worsened the effects of STZ injection suggesting that the BPA exposure affected islet activity in injuring conditions.

Discussion

Epidemiological and experimental studies showed that exposure to BPA elicited alteration in pancreatic islets function and glucose homeostasis. In this work, we studied the effects of BPA on murine primary hepatocytes and pancreatic islets through a toxicogenomic approach, mimicking the worldwide population exposure to BPA low-dose (1 × 10⁻⁹ M). No transcriptional effect was retrieved in primary hepatocytes, even though some effects have been described in other experimental settings, whereas in islets only a few transcripts were inhibited. They were mainly involved in the deregulation of mitochondrial activity and, despite the slight effects detected at mRNA level, the summary of their deregulation of mitochondrial activity and, despite the slight effects detected at mRNA level, the summary of their effect was retrieved in primary hepatocytes, transcriptional effect was retrieved in primary hepatocytes, transcriptional effect was retrieved in primary hepatocytes, whereas the protein induction at longer time of exposure time, whereas the protein induction at longer time, whereas the protein induction at longer time.

The toxicogenomic analysis allowed us to have a snapshot of the global status of the islets, indicating the mitochondrion as the key organelle affected by BPA exposure. Mitochondria are critical for the maintenance of β-cell function coupling glucose stimulus to insulin release and are targets of BPA toxicity. Moreover, alterations in mitochondrial dynamics have been reported among the mechanisms underlying the development of T2D and obesity. Indeed, previous studies indicated that BPA exposure could cause profound mitochondrial structural defects and decrease of mitochondrial activity in pancreatic islets. Here, we report that BPA inhibits the expression of genes involved in mitochondrial activity leading to impairment of the organelle function and its damage. This results in a reduction of insulin secretion following glucose stimulation and in an increase in the apoptosis. Both mechanisms of BPA toxicity are suggested by the toxicogenomic analyses and could contribute to the impairment of blood glucose homeostasis. Other authors report demonstrated that BPA treatment induces an increment in the insulin cellular content in vivo and in vitro after 4 days and 48 h of treatment, respectively, and an increment of insulin release in vitro after 1 h of stimulation. Our results suggest that longer exposure to BPA (48 h) reduces insulin secretion in response to 16 mM glucose. Considering our result, we suggest that the increment of the intracellular insulin content observed by others in the 48 h treated islets is possible owing to the impairment in the secretion mechanism that results in the intracellular accumulation of the protein. Noteworthy, the decrease of the Vapa transcript, codifying for a SNARE protein, here reported could be a part of the mechanism behind this alteration of insulin content/release.

BPA is a xenoestrogen acting through nuclear ERs and membrane-bound ERs. Despite that, the reported bioinformatics analyses predicted the deregulation of other transcription factors, as NF-κB and FOXO families. Therefore, we experimentally confirmed the activation of the NF-κB pathway (Figure 4) and ruled out a clear involvement of the FOXO proteins and, above all, ERα at the analyzed time points. Indeed, previous studies have suggested a direct involvement of ERα, ERβ and GPR30 in mediating low-dose BPA-increased insulin release or protein level. The ERα-dependent increase in the insulin transcript was reported only at shorter exposure time, whereas the protein induction at longer time (48 h) using ERα−/− and ERβ−/− animal. If it is undoubted that ERs have a pivotal role also in insulin release, it must be noted that this effect has been reported at shorter time point and that the activation of other signaling pathways involved in the process has been shown. Therefore, it is not surprising to us that other transcriptional regulators can mediate the reported effects. Although we cannot rule out a role of ERs, as we have specifically inhibited only ERα, we do identify NF-κB pathway as a target of BPA activity in pancreatic islets in our experimental setting. Furthermore, in immortalized thyrocytes, we have already described that ERs are not playing a direct role in mediating BPA transcriptional effects using a specific reporter system underlining meanwhile the role of NF-κB pathway. In light of the reported results, BPA acts in the same way in islet cells activating NF-κB pathway after inducing intracellular ROS level, (Figure 3). These results are in agreement with the findings that BPA exposure increase ROS cellular amount resulting in the activation of the NF-κB pathway in an IKK-dependent manner, here confirmed by the reduction in IkB-α cellular content in treated pancreatic islets.

Figure 5 Effects of BPA on islets functionality. (a) Insulin secretion after 1 h of 16 mM glucose stimulation of islets exposed to 1 × 10⁻⁴ or 1 × 10⁻⁸ M BPA. Data are reported as fold change values. The results are expressed as the mean ± SD of three independent experiments (N = 3). *P-value < 0.05; **P-value < 0.01; ***P-value < 0.001 compared with vehicle-only-treated islets; (b) glycemia measurements in hyperglycemic mice transplanted with pancreatic islets. The animals were intravenously injected (day − 3) with STZ for hyperglycemia induction and transplanted with pancreatic islets from healthy syngenic donors, three days later (day 0). Transplanted islets were cultured for 48 h with (Gr.2, N = 5 and Gr.3, N = 4) or without (control group, Gr.1, N = 5) 1 × 10⁻⁹ M BPA. Gr.2 animals were watered with 50 μg/kg bw/day BPA from the day of STZ injection until the day of killing. *P-value < 0.05; **P-value < 0.01; ***P-value < 0.001 compared with control animals (Gr.1).
Indeed, the inhibited transcripts suggest different mechanisms through which BPA could determine islet failure and death, depicted in Figure 6. The downregulation of transcripts involved in mitochondrial respiratory chain (Uqcrb and Ndufs4) Figure 2a), as well as ATP6v1f (Figure 2b), points to an impaired oxidative phosphorylation. This causes defects in glucose-stimulated ATP production, therefore, directly impairing the insulin secretion. Noteworthy, ATP1b1, a subunit of sodium/potassium-transporting ATPase, is also inhibited in our experiments (Figure 2b) suggesting that alterations in the Ca²⁺ and K⁺ homeostasis could be also involved in islet failure (Figure 6).55 Furthermore, the inhibited expression of respiratory chain genes could promote the leakage of electrons inducing free radical accumulation to which the inhibition of the ROS scavenging transcripts (Sod2 and Gpx3, Figure 2c) could contribute,53 resulting in the NF-κB activation.54 In summary, BPA could exert toxic activity throughout two complementary mechanisms centered on mitochondria: enhancement of the oxidative stress and drop of the ROS scavenging systems, promoting the apoptosis. Their simultaneous impairment amplifies the effects of BPA. Despite the weakness of the detected effects, in terms of gene transcript inhibition, they determine a reduced response of the islets to further injuries, as the exposure to high glucose in vitro and in vivo. Indeed, our findings evidence that the BPA-induced damage cannot be recovered, even after dropping the exposure. In agreement with recent results, where the treatment was performed with a 100 times higher BPA dose,45 we show that BPA exacerbates the diabetogenic effects of STZ. Furthermore, our data suggest that BPA pre-treated islets are impaired and apoptotic, therefore, not efficiently engrafting when transplanted in hyperglycemic mice, even if not longer exposed.

Many evidences show the role of environmental pollutants in diabetes development. Indeed, there were 285 million people worldwide with diabetes in 2010 and this number is estimated to increase by 54% within 2030,54 indicating that factors, other than genetic, could equally contribute. Their clear identification is important to explain the epidemic diffusion of diabetes and to develop preventive measures. To this aim, our findings are significant as they characterize, at transcriptomic level, the reported impairment of islet activity and viability. The toxicogenomic approach reveals different mechanisms through which BPA exerts its effects on pancreatic islets. The future assessment of the real risks related to the BPA exposure needs to turn the attention on the interplay among genetic and multiple environmental factors to better match the real world.

Materials and Methods

Animals. C57/B6 male mice, 22–27 gr., 5–8 weeks old, were obtained from Biogem s.c.a.r.l. (Ariano Irpino, Italy). Mice were housed in polysulfone cages with ad libitum access to water and normal diet under specific pathogen-free facility at 22 °C ± 2° temperature and 55% ± 15% relative humidity with 12 h light: 12 h darkness cycle and 18 ± 2 changes of air per hour. All animal experimentation as respected the regulations and guidelines of Italy and the European Union and the NIH Principles of Laboratory Animal Care (NIH, publication no. 85-23, revised 1985) and have been evaluated and approved by the ethics committee `Comitato Elico per la Sperimentazione Animale’ (CESA) of IGRS, Biogem.

Murine pancreatic islets. Primary cultures of pancreatic islets were prepared by pancreas bile duct perfusion, Collagenase P digestion (0.8 mg/ml Roche GmbH, Mannheim, Germany) and cultured as already described.55 Briefly, islets were purified from exocrine tissue by differential centrifugation on Histopaque 1077 (Sigma-Aldrich, St. Louis, MO, USA) and HBSS (w/o Ca and Mg, Sigma-Aldrich) density gradient. The islet yield and quality was checked on an inverted microscope. The islets were handpicked and cultured at 37 °C with 5% CO₂. Islets were cultured in RPMI 1640 (Sigma-Aldrich)+10% fetal bovine serum (FBS, Sigma-Aldrich)+1% Pen/Strep (Sigma-Aldrich). A total 250–300 islets per 6 cm dish with 3 ml of media were used, avoiding stressing condition. The islets were treated with 1 x ×10⁻⁴M, 1 x ×10⁻⁶M or 1 x ×10⁻⁸M BPA, 25 mM glucose, 3x10⁻⁵ M menadione, 3 mM NAC, 2 x ×10⁻⁵ M BMS-345541, 1x10⁻⁵ M LY294002, 1x10⁻⁵ M IC-182780 or vehicle (DMSO). All the chemicals were purchased from Sigma-Aldrich. Treatments were performed for time points ranging from 3 h to 48 h; at longer time fibroblast proliferation and islets senescence was detected (Supplementary Figure S1). When required, islets were dispersed as previously described.55 For all the experiments, to minimize the effects of mice interindividual variations, the islets from three or more pancreas were pooled and then re-divided for treatments.

Murine primary hepatocytes. Primary culture of murine hepatocytes were obtained by liver collagenase perfusion and digestion as previously described.57 Briefly, after anesthesia and laparotomy, a PE10 catheter was introduced in the portal vein. The liver was perfused with a pre-digestion solution (HBSS w/o Mg and Ca+0.5 mM EGTA+25 mM Hepes+1% Pen/Strep (Sigma-Aldrich), pH 7.4) pre-warmed to 37 °C for 10 min at the rate of 5 ml/min, afterwards perfused with pre-warmed digestion solution (William’s E, Medium+15 mM Hepes+1% Pen/Strep +0.32 mg/ml Collagenase type IV (Sigma-Aldrich), pH 7.4) at the same rate for 10 min. The digested liver was excised and hepatocytes were released by gently shanking into 15 ml of hepatocyte isolation medium (William’s E+4% FBS+1% Pen/Strep (Sigma-Aldrich)). To access the quality of the perfusion, trypan blue staining assessed the cell viability. Then, the cells were plated in collagen type I (Sigma-Aldrich)-coated dishes with hepatocyte isolation medium. After 6 h at 37 °C, the cells were attached and the medium was replaced with hepatocyte growth medium (William’s E+4% heat-inactivated FBS+1% Pen/Strep+50 ng/ml Epidermal Growth Factor+1 mg/ml insulin+10 mg/ml Transferin+1.3 mg/ml Hydrocortisone (Sigma-Aldrich)). The treatments were conducted as described above. Treatments longer than 48 h were not conducted as in vitro mesenchymal transition was observed (Supplementary Figure S1).

MTT and ATP assay. MTT (Sigma-Aldrich) and ATPlite assay (PerkinElmer Life Sciences, Inc, Zaventem, Belgium) were performed on dispersed islets, grown in the appropriate plates, according to manufacturer’s instructions. For MTT, absorbance at 570 nm was read in EnVision 2103 Multilabel Reader (PerkinElmer Life Sciences, Inc). For ATP assay, luminescence was measured in Orion II
microplate luminometer (Berthold Detection Systems GmbH, Pforzheim, Germany). The luminescence level was normalized on protein amount.

RNA extraction and qRT-PCR. RNA was extracted from isolated islets or hepatocytes using Trizol reagent (Life Technologies Italia, Monza, Italy) and quantified with the NanoDrop spectrophotometer ND-1000. For qRT-PCR, 1 μg of total RNA was reverse-transcribed (QuantiTect Reverse Transcription Kit, Qiagen), according to the manufacturer’s instructions. qRT-PCR experiments were conducted using Life Technologies Italia 7900 Real-Time PCR System and Power SYBR Green Master Mix (Life Technologies Italia). The subsequent analysis was performed following Hellymans et al.58 models for qRT-PCR relative quantification and inter-run calibration with proper error. Data were normalized on the relative expression of three reference genes (Gapdh, Tubulin and β-2 microglobulin).59 Primers were designed using NCBI Primer Blast tool (http://www.ncbi.nlm.nih.gov/tools/primer-blast). Their sequences are below:

**Reverse Transcripts (RNA)**

- ATP6v1f: 5′-CTCCTGCTTACATGACCCA-3′ (forward) and 5′-TATGATGCTGCCCGCTTC-3′ (reverse);
- ATPB6f: 5′-ATCAAGAAGGCCTTCTTGCT-3′ (forward) and 5′-ATGCTCTTGGAGGGTACT-3′ (reverse);
- Bcl-x: 5′-CACATGATGAGAAGGAAGGG-3′ (forward) and 5′-CAATGAGGAAGGACAC-3′ (reverse);
- Bcl-2: 5′-ATGACGTGATACCTGACCGGCT-3′ (forward) and 5′-GGGCCCATATGTTCCAAAGGCA-3′ (reverse);
- Bcl-xL: 5′-AACAGCAACGCGGCAAGA-3′ (forward) and 5′-CCGTTACACTCTTCTCCTTCA-3′ (reverse);
- Bcl-2L1: 5′-GCACTTGGCAAATCTCACG-3′ (forward) and 5′-TGAGTGTGGTGC-3′ (forward) and 5′-ATGGATGTTGACTGAGGTTG-3′ (forward);
- IkBα: 5′-GCCTGAGTTCCCTTGAGCAGC-3′ (forward) and 5′-CAATGAGGAAGGACAC-3′ (reverse);
- Caspase-3: 5′-GACGGGTACCAAACTTCCAAC-3′ (forward) and 5′-CCAAGTGCTTCCCCAAGTCAGGA-3′ (forward);
- CACCCGCA-3′ (forward) and 5′-CTCCCACGGACATCTCCTTG-3′ (forward) and 5′-GCTGCGTACGAGGGGCT-3′ (reverse);
- Vpa: 5′-GAGATTGTGTTTGGAAACGCGA-3′ (forward) and 5′-GGTCCGCTTTGTTGGTGGAGG-3′ (forward);
- Pdcdg: 5′-GCCGGGCGTCTGTTGC-3′ (forward) and 5′-GCTCTGATCGAGGGGCT-3′ (reverse);
- H2A-Z: 5′-GACGTGACGGCTGATG-3′ (forward) and 5′-AAGACGATCTGCAGCTGAG-3′ (forward);
- Brm: 5′-GCAACGACAGACTGCTGACT-3′ (forward) and 5′-TGTCGGTTCTCTGGTGC-3′ (forward);
- Zn-finger: 5′-ACCCGTGAGTCCGCTGATG-3′ (forward) and 5′-AACAGGTCTCAGGCAAGGCA-3′ (forward);
- Brm: 5′-GCAACGACAGACTGCTGACT-3′ (forward) and 5′-TGTCGGTTCTCTGGTGC-3′ (forward);
- Tnfα: 5′-ACCAGTGTGCTGCTGATG-3′ (forward) and 5′-CACCCACGGCTGTGAGGCTC-3′ (reverse);
- Tuba: 5′-CAACACCTTCTCTGAGGAGC-3′ (forward) and 5′-CATGACCTTCTGCTCAGGATG-3′ (forward).

**Microarray.** Gene expression profiling experiments and data analyses were conducted as elsewhere described.59 The Affymetrix platform was used. cRNA was generated by using the Affymetrix One-Color Target Labeling and Control Reagent kit (Affymetrix Inc., Foster City, CA, USA), according to the manufacturer’s instructions, starting from 5 μg of total RNA. Biotinylated cRNA was hybridized to the Mouse MOE 430 2.0 Genome Arrays (Affymetrix Inc.). Chips were washed and dried, and the luminescence level was normalized on protein amount. Equal protein amount (20 μg) from the islets lysates were separated by SDS-PAGE. Membranes were probed with primary antibody against IkBα (sc-371, Santa Cruz Biotechnology Inc., Dallas, TX, USA). Mouse β-actin antibody (Sigma-Aldrich) was used to normalize protein levels. Band intensities were quantified by ImageJ software (1.48v, National Health Institute, NIH, Bethesda, MD, USA).

**Mitochondrial membrane potential measurements.** Mitochondrial JC-1 assay kit (Life Technologies Italia) was used to measure mitochondrial membrane potential. The assay was performed on dispersed islets, grown in the appropriate plates, according to manufacturer’s instructions. Fluorescence was measured with EnSpire Multimode Plate Reader (ex/em. 514/530, 575/590 nm, PerkinElmer Life Sciences, Inc) after treatment with BPA 1 × 10−5 M at different times. Treatment with 2 × 10−6 M carbonyl cyanide 3-chlorophenylhydrazide (CCCP) for 5 min was used as positive control. The fluorescence level was normalized on protein amount. A decrease in the red/green fluorescence intensity ratio was taken as indicator of mitochondrial depolarization.

**Fluorescence levels were measured bi-weekly until killing at day 15 post transplant.**
Statistical and bioinformatics analyses. The statistical analyses have been performed with Student's t-test, unless otherwise indicated. In all the cases, probability P-values below 0.05 were considered significant. * , **, *** indicate P-value < 0.05, < 0.01, < 0.001, respectively. Data from at least three independent experiments were considered for the statistical analysis. Unless otherwise indicated, data are reported as fold change values calculated as ratio between average results in treated and control samples. The results are expressed as the mean ± SD of three independent experiments. Hypothetical transcriptional factors binding sites in the gene promoter were identified loading their sequence ranging from −1000/−50 bp in Genomatix suite software (Genomatix Software GmbH, http://www.genomatix.de), choosing a relative profile score of 80%.

Conflict of Interest
The authors declare no conflict of interest.

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