Formoterol protects against platelet-activating factor-induced effects in asthma

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ABSTRACT: Platelet-activating factor (PAF) is an inflammatory mediator that provokes neutropaenia, bronchoconstriction and gas exchange defects due to exudation of bulk plasma within the airways. While the inhibitory effects of short-acting β₂-agonists on PAF-induced disturbances have been consistently shown, those of long-acting β₂-agonists are less convincing.

To further explore the mechanisms involved in PAF challenge in asthma, 12 patients (forced expiratory volume in one second, 90±4% predicted) were investigated 2 h after inhaled formoterol (18 μg), in a double-blind, placebo-controlled, crossover design following PAF (18 μg) inhalation.

Compared with the placebo, at 5 min, premedication with formoterol reduced PAF-induced cough and dyspnoea, and attenuated increased respiratory system resistance (by 67%) and arterial deoxygenation (by 50%). Likewise, ventilation-perfusion (V′/Q′) inequality improved, as reflected by the dispersion of pulmonary blood flow (by 63%) and an overall index of V′/Q′ heterogeneity (by 71%). In contrast, PAF-induced facial flushing, neutropaenia and subsequent rebound neutrophilia remained unchanged.

The improvement in gas exchange abnormalities shown after platelet-activating factor in patients with asthma pretreated with formoterol at the recommended clinical dose may reflect, in addition to its class effects, an anti-exudative effect of formoterol in the airways.

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Platelet-activating factor (PAF) is a potent ether-linked phospholipid mediator of inflammation, which has been suggested to play a pathogenic role in bronchial asthma [1]. PAF induces neutropaenia, bronchoconstriction and gas exchange defects due to exudation of bulk plasma within the airways in both normal subjects and asthmatics [2]. These gas exchange abnormalities include the development of areas with low ventilation-perfusion (V′/Q′) units identical to those shown in patients with spontaneous acute asthma [2–4]. PAF potentiates its effects by generating secondary release of other inflammatory mediators, such as leukotrienes (LT)s, via the activation of phospholipase A₂ [5, 6].

Both inhaled short-acting and long-acting β₂-agonists (LABAs) remain one of the mainstays of asthma therapy. In addition to their potent bronchodilator effects, these agents also exhibit nonbronchodilator properties, such as anti-exudative efficacy [7, 8]. The current authors have previously shown that inhalation of salbutamol (300 μg) is able to suppress all PAF-induced systemic, cellular and functional abnormalities in normal subjects [9] and asthmatics [10]. However, the LABA salmeterol (50 μg, b.i.d.) administered for 1 week failed to inhibit neutrophil sequestration and bronchoconstriction caused by PAF [11]. In contrast, the rapid-onset LABA formoterol demonstrated potent anti-exudative and bronchodilator properties in healthy individuals [7] and in animal models [12]. Therefore, the authors of this paper investigated the effects of a therapeutic dose of inhaled formoterol fumarate in patients with mild asthma and further explored the systemic, cellular, lung mechanical and gas exchange alterations provoked by PAF inhalation.

Materials and methods

Subjects

Twelve nonsmoking, mild asthmatics (eight males; seven atopics) were recruited for the study (table 1) after approval by the Ethical Committee of Hospital Clinic, Barcelona. All patients gave informed written consent after the aims, risks and potential benefits of the study were explained to them. Inclusion criteria were: lack of asthma exacerbation within the preceding 6 weeks; forced expiratory volume in one second (FEV₁) >70% predicted; positive methacholine bronchial challenge (provocation dose causing a 20% fall in FEV₁ <1.9 μmol); positive PAF challenge (20% increase of baseline...
Platelet-activating factor challenge

A randomised, double-blinded, placebo-controlled, two-period crossover design was used. All patients were challenged on two occasions, 1 week apart, with inhaled PAF 2 h after administration of placebo and formoterol. During the challenge the patients breathed room air and were seated in a semirecumbent position. All asthma medication was withheld 24 h before the arrival to the laboratory. After the establishment of adequate steady-state conditions, a first set of duplicate measurements was taken (B0). Maintenance of steady-state conditions after PAF challenge was demonstrated by stability (±5%) of ventilatory and haemodynamic outcomes, and by the close agreement between duplicate measurements of mixed expired and arterial O2 and CO2 (within ±5%). These conditions were present in all patients throughout the entire period of study. Accordingly, the residual sum of squares (RSS) [13], a reliable descriptor of the goodness of the fit of inert gas data, was 4.4±0.3 for all MIGET studies following PAF challenge (RSS <10.6 in 94% of sets) [14]. A second set of measurements was made 2 h after placebo/formoterol administration (B1), and the patient was then challenged with PAF (C16) (1-0-hexadecyl-2-acetylsn-glycero-3-phosphocholine) (18 µg) (Novabiochem AG, Laufelfingen, Switzerland). Preparation of the PAF solution and details of the PAF challenge have been previously reported in full [3, 4]. Duplicate measurements were then made at 5, 15, and 45 min after PAF inhalation. All sets of measurements consisted of the following steps in sequence: ventilatory recordings; respiratory and inert gas (mixed venous and arterial) and circulating white blood cells; and haemodynamic and Rs measurements.

Analysis of data

Results are expressed as either the arithmetic mean±SEM or 95% confidence interval. Comparison of baseline conditions before and 2 h after placebo/formoterol and before PAF challenge, and both the effects of PAF challenge and following administration of vehicle/formoterol were assessed using a two-way repeated measures analysis of variance (two-way analysis of variance). Whenever an interaction was found between the effects of PAF challenge and those shown after administration of the two pretreatments, differences between placebo and formoterol at each time point were assessed with a post-hoc paired t-test. A Chi-squared test was used for noncategorical variables (symptoms). Significance was set at p<0.05 in all instances.

Results

Baseline findings (before platelet-activating factor)

Anthropometric and functional data were within normal limits and close to those reported in previous studies [15].

| Table 1. – Patient anthropometric data |
|---------------------------------------|
| Age yrs                               | 24±1 |
| Height cm                             | 170±2 |
| Weight kg                             | 72±3 |
| FEV1/L                                | 3.6±0.2 |
| FEV1 % pred                           | 90±54 |
| PD20 normal <1.9 µmol                 | 0.7±0.1 |

Data are presented as mean±SEM. FEV1: forced expiratory volume in one second; % pred: % predicted; PD20: provocative dose of methacholine causing FEV1 to fall 20% from baseline.

Respiratory system resistance (Rrs) 5 min after PAF (18 µg); maintenance therapy with short-acting β2-adrenergics (SABAs) and/or inhaled corticosteroids; no previous treatment with systemic steroids; and absence of any systemic or cardiopulmonary disease other than asthma. Maintenance therapy included rescue medication with SABAs alone (four patients) or combined with either regular (four patients) or seasonal inhaled glucocorticoid treatment (three patients); the remaining patient added an oral leukotriene receptor antagonist to seasonal inhaled glucocorticoids and SABAs on clinical demand.

Measurements

Rrs was measured by the forced oscillation technique and its analysis restricted to 8 Hz [3, 4]. Both minute ventilation and respiratory rate were measured using a calibrated Wright spirometer (Respirometer MK8; BOC-Medical, Essex, UK). A three-lead electrocardiogram, cardiac frequency, and systemic pressure were continuously recorded throughout the whole study (HP 7830A Monitor and HP 7754B Recorder; Hewlett-Packard, Waltham, MA, USA). Both oxygen (O2) uptake and carbon dioxide (CO2) production were calculated from mixed expired O2 with a Zirconia analyser (MCG Graphics Corporation, St. Paul, MN, USA) and CO2 concentration was measured via a nondispersive infrared analyser (Model CPX/D; MCG Medical Graphics Corporation). Blood samples were collected under anaerobic conditions through a catheter inserted into the radial artery. Arterial O2 tension (PaO2), arterial CO2 tension and pH were analysed in duplicate using standard electrodes, and haemoglobin concentration was measured using a co-oximeter (Ciba corning AstraZeneca, Madrid, Spain; 9 mol 0.7 µmol 0.2 mg). The patients were given allocation numbers prepared in blocks of four. Blindness was maintained by identical appearance of active and vehicle administrations.

A randomised, double-blinded, placebo-controlled, two-period crossover design was used. All patients were challenged on two occasions, 1 week apart, with inhaled PAF 2 h after administration of placebo and formoterol. During the challenge the patients breathed room air and were seated in a semirecumbent position. All asthma medication was withheld 24 h before the arrival to the laboratory. After the establishment of adequate steady-state conditions, a first set of duplicate measurements was taken (B0). Maintenance of steady-state conditions after PAF challenge was demonstrated by stability (±5%) of ventilatory and haemodynamic outcomes, and by the close agreement between duplicate measurements of mixed expired and arterial O2 and CO2 (within ±5%). These conditions were present in all patients throughout the entire period of study. Accordingly, the residual sum of squares (RSS) [13], a reliable descriptor of the goodness of the fit of inert gas data, was 4.4±0.3 for all MIGET studies following PAF challenge (RSS <10.6 in 94% of sets) [14]. A second set of measurements was made 2 h after placebo/formoterol administration (B1), and the patient was then challenged with PAF (C16) (1-0-hexadecyl-2-acetylsn-glycero-3-phosphocholine) (18 µg) (Novabiochem AG, Laufelfingen, Switzerland). Preparation of the PAF solution and details of the PAF challenge have been previously reported in full [3, 4]. Duplicate measurements were then made at 5, 15, and 45 min after PAF inhalation. All sets of measurements consisted of the following steps in sequence: ventilatory recordings; respiratory and inert gas (mixed venous and arterial) and circulating white blood cells; and haemodynamic and Rs measurements.

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Results

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after PAF exposure (table 3 and fig. 1). Compared with formoterol, at 5 min, $R_s$ (p<0.003) and $P_AaO_2$ (p<0.05) increased and $P_aO_2$ (p<0.05) fell moderately to severely. Arterial deoxygenation was paralleled by the development of moderate-to-severe ventilation/perfusion ratio ($V'AV/Q'$) inequality, as demonstrated by abnormal increases in two of its best descriptors: the dispersion of pulmonary blood flow (LogSD Q) (p<0.02) and an overall index of $V'AV/Q'$ heterogeneity (DISP R–E*) (p<0.005). At 15 min, $R_s$ (p<0.0003), $P_aO_2$ and $P_AaO_2$ (p<0.01 each) remained considerably altered and $V'AV/Q'$ imbalance persisted (LogSD Q, p<0.02; DISP R–E*, p<0.001). By 45 min, $R_s$ (p<0.004), $P_AaO_2$ (p<0.03), LogSD Q (p<0.002), and DISP R–E* (p<0.01) were still mildly increased and $P_aO_2$ was slightly reduced (p<0.05).

In contrast, circulating blood neutrophils and ventilatory and haemodynamic parameters, and all the other gas exchange indices, including arterial pH (at baseline all were within normal limits), remained stable. No patient needed rescue medication.

**Effects of platelet-activating factor after placebo**

All patients noticed facial flushing and feeling of warmth, nine felt shortness of breath, and six coughed immediately after PAF exposure (table 3 and fig. 1). Compared with formoterol, at 5 min, $R_s$ (p<0.003) and $P_AaO_2$ (p<0.05) increased and $P_aO_2$ (p<0.05) fell moderately to severely. Arterial deoxygenation was paralleled by the development of moderate-to-severe ventilation/perfusion ratio ($V'AV/Q'$) inequality, as demonstrated by abnormal increases in two of its best descriptors: the dispersion of pulmonary blood flow (LogSD Q) (p<0.02) and an overall index of $V'AV/Q'$ heterogeneity (DISP R–E*) (p<0.005). At 15 min, $R_s$ (p<0.0003), $P_aO_2$ and $P_AaO_2$ (p<0.01 each) remained considerably altered and $V'AV/Q'$ imbalance persisted (LogSD Q, p<0.02; DISP R–E*, p<0.001). By 45 min, $R_s$ (p<0.004), $P_AaO_2$ (p<0.03), LogSD Q (p<0.002), and DISP R–E* (p<0.01) were still mildly increased and $P_aO_2$ was slightly reduced (p<0.05). In contrast, circulating blood neutrophils and ventilatory and haemodynamic parameters, and all the other gas exchange indices, including arterial pH (at baseline all were within normal limits), remained stable. No patient needed rescue medication.

**Effects of platelet-activating factor after formoterol**

Compared to the effect of vehicle, only three patients had dyspnoea (p<0.02) and one patient had cough (p<0.03) after PAF; in contrast, facial flushing remained almost unchanged.

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**Table 2. – Baseline data for placebo and formoterol studies**

|                  | Placebo       | Formoterol    |
|------------------|---------------|---------------|
| Neutrophils $\times 10^9$ L$^{-1}$ | 3.3±0.3       | 3.3±0.4       |
| $R_s$ cm H$_2$O L$^{-1}$s$^{-1}$   | 6.1±0.5       | 5.3±0.4       |
| $P_aO_2$ mmHg     | 102.4±3.2     | 100.3±2.2     |
| $P_AaO_2$ mmHg    | 2.4±1.6       | 2.7±1.7       |
| LogSD Q           | 0.48±0.05     | 0.46±0.04     |
| LogSD V           | 0.45±0.03     | 0.44±0.03     |
| DISP R–E*         | 3.24±0.48     | 2.98±0.41     |

Data are presented as mean±SEM. $R_s$: resistance of respiratory system; $P_aO_2$: arterial oxygen tension; $P_AaO_2$: alveolar–arterial oxygen tension difference; LogSD Q: dispersion of pulmonary blood flow distribution; LogSD V: dispersion of alveolar ventilation distribution; DISP R–E*: retention minus excretion of inert gases corrected for deadspace. mmHg×0.133=kPa.

There were no differences between the placebo and the formoterol subjects (table 2 and fig. 1).

**Fig. 1. –** Time course of a) respiratory system resistance, b) an overall index of ventilation-perfusion mismatching (expressed as an overall index of ventilation/perfusion ratio heterogeneity (DISP R–E*)), c) peripheral blood neutrophils, and d) arterial oxygen tension before and after platelet-activating factor challenge. Data are presented as mean±SEM. mmHg×0.133=kPa. BO: baseline measurements before pretreatment; B1: measurements 2 h after pretreatment, before platelet-activating factor exposure. Closed arrows show placebo/formoterol administration, open arrows show platelet-activating factor challenge. ●: formoterol; ○: placebo.
with no response in two patients (table 3 and fig. 1). At 5 min, PAF-induced increases in Rs (by 67%) and PAaO2 (by 53%) and hypoxaemia (by 50%) were substantially attenuated; similarly both abnormal LogSD Q (by 63%) and DISP R-E* (by 71%) were largely ameliorated. By 15 min, Rs (by 85%), PaO2 (by 58%), PaAaO2 (by 71%), and P/AO2 inequality continued to improve and all gas exchange markers were already within normal limits. At 45 min, all these outcomes were normalised. Abnormal neutrophil kinetics remained unaltered throughout the study.

**Discussion**

The novel finding of the current study was that, in addition to inhibiting bronchoconstriction, a clinically recommended dose of inhaled formoterol used in patients with mild asthma considerably protected against arterial desoxygenation and underlying ventilation-perfusion imbalance after PAF. In contrast, formoterol was unable to modulate either facial flushing or neutrophil kinetics. Based on previous work by the authors in both normal subjects [3, 8, 9] and mild asthmatics [4, 10, 15], the authors postulate that PAF-induced pulmonary blood flow distribution; DISP R-E: retention minus excretion inert gases corrected for deadspace; NS: not significant.

**Table 3. – Data after placebo (P) and formoterol (F) pretreatments on each platelet-activating factor challenge**

| Parameter | Baseline | 5 min | 15 min | 45 min | p-value<sup>ab</sup> |
|-----------|----------|-------|--------|--------|----------------------|
| Neutrophils ×10<sup>9</sup>·L<sup>-1</sup> | P 2.9 (2.3–3.5) | 1.1 (0.5–1.6) | 5.4 (4.6–6.3) | 5.6 (4.6–6.6) | NS |
| | F 3.1 (2.6–3.7) | 1.6 (0.9–2.4) | 4.6 (3.0–6.2) | 4.9 (3.2–6.6) | |
| Rs cm H<sub>2</sub>O·L<sup>-1</sup>·s<sup>-1</sup> | P 6.2 (4.9–7.4) | 17.0 (13.0–21.0) | 14.1 (10.5–17.7) | 10.1 (7.0–13.1) | 0.0001 |
| | F 4.7 (3.7–5.7) | 8.3* (5.2–11.4) | 6.9* (4.4–7.6) | 5.1* (4.2–5.9) | |
| PAoO<sub>2</sub> mmHg | P 103.1 (98.2–108.0) | 74.9 (65.4–84.5) | 82.5 (74.2–90.9) | 96.0 (89.8–102.2) | 0.02 |
| | F 104.5 (98.4–110.6) | 90.4* (81.1–99.8) | 97.3* (89.9–104.6) | 103.8* (98.1–109.6) | |
| PAaO<sub>2</sub> mmHg | P 1.8 (-0.1–4.3) | 27.7 (18.2–37.3) | 18.5 (8.9–28.2) | 5.8 (1.1–16.8) | 0.009 |
| | F 1.5 (-0.7–3.0) | 11.9* (1.3–22.4) | 5.9* (0.1–11.7) | 1.7 (-0.1–4.1) | |
| LogSD Q | P 0.49 (0.40–0.57) | 1.01 (0.80–1.21) | 0.80 (0.64–1.00) | 0.63 (0.51–0.76) | 0.02 |
| | F 0.41 (0.33–0.48) | 0.59* (0.37–0.82) | 0.25* (0.13–0.69) | 0.34* (0.26–0.52) | |
| DISP R-E<sup>ab</sup> | P 3.31 (2.50–4.12) | 9.38 (7.01–11.74) | 5.95 (4.17–7.74) | 4.14 (2.96–5.32) | |
| | F 4.14 (2.96–5.32) | 4.06 (3.09–5.03) | 3.63 (2.83–4.43) | 2.80 (2.02–3.58) | |

Data are presented as mean (95% confidence interval) unless otherwise stated. All data correspond to 12 patients, except for neutrophil and inert gas variables (n=11). mmHg=0.133=kPa. Rs: resistance of respiratory system; PaO2: arterial oxygen tension; PAaO2: alveolar-arterial oxygen tension difference; logSD Q: dispersion of pulmonary blood flow distribution; DISP R-E: retention minus excretion inert gases corrected for deadspace; NS: not significant. <sup>a</sup>: Significance of the interaction between the effects of platelet-activating factor challenge and pretreatment with placebo and formoterol using a two-way repeated measures analysis of variance. *: p<0.05 for comparison with placebo.
previous suggestions [7] that a vascular antipermeability property can be an important step of its mechanisms of therapeutic effect of long-acting $\beta_2$ agonists in asthma. Anti-exudative properties of long-acting $\beta_2$ agonists may enhance their antiasthma efficacy beyond bronchodilation and add potential for reducing asthma exacerbations [30].

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