Abstract. In the present study, rats were exposed from the 8th week after birth and for the ensuing 8 weeks to diets containing either starch or fructose (64% w/w) and sunflower oil (5%). Two further groups of rats were exposed to the fructose-containing diet with substitution of part (1.6%) of the sunflower diet by an equal amount of either salmon oil rich in long-chain polyunsaturated ω3 fatty acids or safflower oil reach in long-chain polyunsaturated ω6 fatty acids. The insulin content of the islets and their secretory response to D-glucose (5.6, 8.3 and 16.7 mM), to the combination of D-glucose (5.6 mM) and D-fructose (10.0 mM) and to 2-ketoisocaproate (10.0 mM) were then measured. In the sunflower oil-fed rats, the dietary substitution of starch by fructose decreased basal insulin output, lowered the apparent Km for the insulinotropic action of D-glucose and altered the insulinotropic efficiency of the latter hexose relative to that of other nutrients. Some of these secretory perturbations were opposed by the enrichment of the diet in long-chain polyunsaturated fatty acids, especially ω3 fatty acids. It is proposed that these changes in B-cell secretory behaviour may be linked, in part at least, to both the apparent caloric efficiency of each diet, and hence to the regulation of the islet content in endogenous nutrients, and to alteration of insulin sensitivity considered as a major feature of the present animal model of metabolic syndrome.

Introduction
In the two preceding reports in this series (1,2), attention was mainly drawn to the perturbation of glucose homeostasis found in rats exposed to diets containing sunflower oil and either starch or D-fructose as the source of carbohydrates and in fructose-fed rats exposed to diets enriched in either ω3 fatty acids rats or C18:2ω6 rats. The present report deals with the insulin secretory behaviour of pancreatic islets isolated from the same four groups of rats.

Materials and methods
The four groups of rats examined in this study were defined in the second report in this series (1). The methods used to measure insulin release over 90 min incubation at 37˚C and the final insulin content of the islets were previously described (3). Briefly, groups of 8 islets each were incubated for 90 min at 37˚C in 1.0 ml of a salt-balanced medium (4) containing 0.5 mg/ml bovine serum albumin and equilibrated against a mixture of O2/CO2 (95/5, v/v). After incubation, the islets were examined for their insulin content. The insulin and protein content (5) of freshly isolated islets was also measured. All results are expressed as mean values (± SEM) together with the number of individual determinations (n) or degree of freedom (df). The statistical significance of difference between mean values was assessed by use of Student's t test.

Results

Insulin release recorded in the presence of 5.6 mM D-glucose.
As documented in Table I, the close-to-basal release of insulin recorded over 90 min incubation in the presence of 5.6 mM D-glucose was somewhat higher (p<0.04) in the (Ssun) rats than in the 3 groups of fructose-fed animals. Pooling together all available data, it averaged 51.4±4.2 µU/islet per 90 min (n=67).

Insulin release recorded in the presence of both 5.6 mM D-glucose and 10.0 mM D-fructose. The increment in insulin output caused by 10.0 mM D-fructose in islets also exposed to 5.6 mM D-glucose was highly significant in all cases (p<0.005 or less), whether expressed in absolute or relative terms (Table I). In absolute terms, such an increment was significantly lower (p<0.02 or less) in the fructose-exposed
Table I. Insulin output.

<table>
<thead>
<tr>
<th>Rats</th>
<th>Ssun</th>
<th>Fsun</th>
<th>Fsal</th>
<th>Fsaf</th>
</tr>
</thead>
<tbody>
<tr>
<td>D-glucose 5.6 mM</td>
<td>77.8±12.9 (16)</td>
<td>35.6±6.1 (16)</td>
<td>46.7±5.9 (18)</td>
<td>46.5±3.7 (17)</td>
</tr>
<tr>
<td>D-glucose 5.6 mM + D-fructose 10.0 mM</td>
<td>170.8±23.9 (20)</td>
<td>75.5±6.4 (15)</td>
<td>117.6±20.4 (18)</td>
<td>77.9±10.5 (17)</td>
</tr>
<tr>
<td>D-glucose 8.3 mM</td>
<td>91.8±9.1 (19)</td>
<td>58.8±9.1 (17)</td>
<td>71.8±7.0 (18)</td>
<td>59.3±4.7 (17)</td>
</tr>
<tr>
<td>D-glucose 16.7 mM</td>
<td>185.2±15.4 (20)</td>
<td>101.2±13.5 (17)</td>
<td>170.5±21.7 (19)</td>
<td>147.6±18.4 (16)</td>
</tr>
<tr>
<td>2-ketoisocaproate 10.0 mM</td>
<td>83.5±9.4 (20)</td>
<td>60.1±10.4 (17)</td>
<td>68.4±8.3 (19)</td>
<td>54.5±3.9 (17)</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Increment in insulin output (µU/islet per 90 min)</th>
<th>Ssun</th>
<th>Fsun</th>
<th>Fsal</th>
<th>Fsaf</th>
</tr>
</thead>
<tbody>
<tr>
<td>∆ 10.0 mM D-fructose</td>
<td>92.5±16.9 (30)</td>
<td>38.3±11.4 (25)</td>
<td>65.5±16.9 (30)</td>
<td>31.5±10.6 (28)</td>
</tr>
<tr>
<td>∆ 8.3 mM D-glucose</td>
<td>11.3±7.6 (29)</td>
<td>24.4±6.7 (27)</td>
<td>24.6±5.9 (30)</td>
<td>12.7±5.9 (28)</td>
</tr>
<tr>
<td>∆ 16.7 mM D-glucose</td>
<td>103.1±12.2 (30)</td>
<td>66.9±12.4 (27)</td>
<td>121.0±19.1 (31)</td>
<td>99.7±13.0 (27)</td>
</tr>
<tr>
<td>∆ 10.0 mM 2-ketoisocaproate</td>
<td>1.8±8.2 (30)</td>
<td>25.2±8.8 (27)</td>
<td>19.6±6.6 (31)</td>
<td>7.9±5.2 (28)</td>
</tr>
</tbody>
</table>

*The increments in insulin output refer to the difference found in each experiment above the mean corresponding value recorded in the presence of 5.6 mM D-glucose.

rats than in the (Ssun) rats, whether the diet of the former rats was enriched or not with ω6 fatty acid. As a matter of fact, there was no significant difference (p>0.67) between these two groups of rats, with an overall mean value for the D-fructose-induced increment in insulin output of 34.7±7.7 µU/islet per 90 min (df=53) as distinct (p<0.001) from 92.5±16.9 µU/islet per 90 min (df=30) in the control animals. In the (Fsal) rats, the D-fructose-induced increment in insulin output averaged 65.5±16.9 µU/islet per 90 min (df=30). It was thus higher albeit not quite significantly so (p<0.07), than that recorded in the (Fsun) or (Fsaf) rats and no more significantly different (p>0.26) from the D-fructose-induced increment in insulin output recorded in the (Ssun) rats.

Essentially comparable information was reached when considering the relative values for the D-fructose-induced increment in insulin output (Table I). Briefly, such relative increment yielded mean values comparable in (Ssun) rats (112.3±20.7%; df=30) and (Fsal) rats (125.4±2.4%; df=30), and lower values in the (Fsun) rats (89.0±25.6%; df=25) or (Fsaf) rats (67.0±21.4%; df=28). The difference between the mean values found in the former 2 groups of rats (118.8±15.8%; df=60) and latter 2 groups of rats (77.4±16.5%; df=53) was close to achieving statistical significance (p<0.08).

Insulin release recorded in the presence of 16.7 mM D-glucose. A somewhat different situation prevailed when considering the increment in insulin output evoked by a rise in D-glucose concentration from 5.6 to 16.7 mM. In absolute terms, the D-glucose-induced increment in insulin output in the (Fsal) rats remained lower (p<0.05 or less) than that found in either the (Ssun) or (Fsal) rats, and even became lower (p<0.06) from that recorded in the (Fsaf) rats. In relative terms, however, the lowest value was recorded in the (Ssun) rats (197.3±36.9%; df=30), it being nevertheless not significantly different (p>0.35) from the mean value found in the other 3 groups of rats (231.5±19.8%; df=85).

The difference between the diet-induced changes in the secretory response to D-fructose, on one hand, and a rise in D-glucose concentration from 5.6 to 16.7 mM, in the other hand, was best documented by the ratio between corresponding variables in these two instances. Indeed, in this respect, comparable results were obtained for i) the ratio between the mean values for the absolute increments in insulin output, ii) the ratio between the mean values for the relative increments in insulin release, and iii) the paired ratio in each individual experiment between the increments in insulin secretion. As illustrated in Fig. 1, the ratio between the secretory response to the rise in D-glucose concentration and that to D-fructose yielded, as judged by these three criteria, higher mean values in the 3 groups of fructose-exposed rats than in (Ssun) animals. More precisely, the concerned ratio, when compared to that recorded in the (Ssun) animals, was not significantly higher (p<0.1) in the (Fsal) rats, close to being significantly higher (p<0.06) in the (Fsun) rats, and obviously higher (p<0.005) in the (Fsaf) rats. No significant difference (p>0.16) was observed between the latter two groups of rats with an overall mean value (expressed...
by reference to the control animals) amounting to 191.4±26.2%, as distinct (p<0.04) from only 136.1±17.4% in the (Fsal) rats.

The diet-induced changes in the secretory response to 16.7 mM D-glucose, as distinct from the combination of 5.6 mM D-glucose and 10.0 mM D-fructose, were also documented by comparing the absolute values for insulin output, the ratio between the mean values for the relative increments in insulin output, the paired ratio in each individual experiment between the increments in insulin secretion, and the overall mean value derived from the former three ratios (df=87-90), the mean measurements collected in the (Ssun) rats being in all cases taken as the unity reference value.

Figure 1. Ratio between the increments in insulin output above the close-to-basal value recorded at 5.6 mM D-glucose caused either by a rise in D-glucose concentration to 16.7 mM or by the incorporation of 10.0 mM D-fructose in the incubation medium in (Ssun) rats (vertically hatched columns), (Fsal) rats (obliquely hatched columns), (Fsaf) rats (open columns) or (Fsun) rats (horizontally hatched columns). Mean values (± SEM; df=27-31) refer from left to right to the ratio between the mean values for the absolute increments in insulin output, the paired ratio in each individual experiment between the increments in insulin secretion, and the overall mean value derived from the former three ratios (df=87-90), the mean measurements collected in the (Ssun) rats being in all cases taken as the unity reference value.

Figure 2. Increments in insulin output evoked by a rise in D-glucose concentration from 5.6 to 8.3 mM, expressed relative to those provoked by a rise in D-glucose concentration from 5.6 to 16.7 mM. Same presentation as in Fig. 1.

Insulin release recorded in the presence of 8.3 mM D-glucose. Whether expressed in absolute or relative terms, the increase in insulin output evoked by a rise in D-glucose concentration from 5.6 to 8.3 mM failed to differ significantly in the 4 groups of rats considered in the present study, with overall mean values (df=14) of 18.2±3.3 µU/islet per 90 min (p<0.001) and 50.3±7.5% (p<0.001). The latter relative increment achieved statistical significance (p<0.03 or less) in all 4 groups of rats.

When the increments in insulin output evoked by a rise in D-glucose concentration from 5.6 mM to 8.3 mM were expressed relative to those provoked by a rise in the D-glucose concentration from 5.6 to 16.7 mM, a comparable hierarchy between the 4 groups of rats was observed, whether the results under consideration referred to the mean absolute increments in insulin output, mean relative increments in insulin output, or paired ratio in each individual experiment between the increments in insulin output under consideration. Thus, as illustrated in Fig. 2, the overall mean values derived from these three criteria averaged 14.6±3.8% (df=77) in the (Ssun) animals, 13.4±3.5% (df=73) in the (Fsaf) rats, 22.8±3.2% (df=78) in the (Fsal) rats and 33.9±5.4% (df=71) in the (Fsun) rats. The highly significant difference (p<0.005) between the first and last of these mean percentages suggests that chronic exposure to a fructose-rich diet lowered the apparent Km for the insulin secretory response of the islets to increasing concentrations of extracellular D-glucose. Incidentally, in this comparison between the secretory rates recorded at 8.3 and 16.7 mM D-glucose, the overall mean reference values averaged 100.0±8.6% (df=80) in the (Ssun) rats, 100.0±8.3% (df=71) in the (Fsaf) rats, 100.0±7.9% (df=81) in the (Fsal) rats, and 100.0±10.6% (df=71) in the (Fsun) rats.

Insulin release recorded in the presence of 10.0 mM 2-ketoisocaprate. The secretion of insulin provoked by 10.0 mM
2-ketoisocaproate, when compared to that recorded within the same experiments in the presence of 5.6 mM D-glucose, yielded mean increments in insulin output not significant different from one another in the 4 groups of rats, with an overall mean values of 13.5±3.7 µU/islet per 90 min (df=116; p<0.001). In this case, the relative increments in insulin output averaged 36.9±8.2% (df=116; p<0.001).

Since the basal release of insulin prevailing in the absence of any exogenous nutrient was not measured in the present experiments, the absolute values for insulin output recorded in the presence of 2-ketoisocaproate (10.0 mM), as measured in the absence of D-glucose, were compared to those found in the presence of 8.3 or 16.7 mM D-glucose.

The absolute values for insulin output recorded in the presence of 2-ketoisocaproate (10.0 mM) were slightly lower (p<0.02) in the fructose-fed rats (61.3±4.6 µU/islet per 90 min; n=53) than in the (Ssun) rats (83.5±9.4 µU/islet per 90 min; n=20). Such was also the case in the islets exposed to 8.3 mM D-glucose with mean values of 91.8±9.1 µU/islet per 90 min; n=19) in (Ssun) animals, as compared (p<0.003) to only 63.5±4.1 µU/islet per 90 min (n=52) in the fructose-fed rats. As a matter of fact, the absolute values for insulin output in the presence of 2-ketoisocaproate averaged 94.9±7.2% (n=73; p>0.57) of the mean corresponding values found in the same group of rats in islets exposed to 8.3 mM D-glucose (100.0±5.4%; n=71). As expected from these findings, the difference found in each experiment (n=3) and in each type of rat between the mean secretory rates recorded in the presence of 2-ketoisocaproate, tested in the absence of D-glucose, and in the presence of only 5.6 mM D-glucose, provided a highly significant (p<0.001) overall positive average (+ 13.5±3.7 µU/islet per 90 min; df=116). The absolute values for insulin output recorded in the presence of 2-ketoisocaproate, when compared to those found in the presence of 16.7 mM D-glucose also failed to differ significantly (p>0.29 or more) in the four groups of rats, with an overall mean values of 45.2±3.2% (n=73) relative to the corresponding reference measurements made in the presence of 16.7 mM D-glucose (100.0±5.7%; n=72).

In order to compare the secretory response to 2-ketoisocaproate and 16.7 mM D-glucose, the difference in the absolute values for insulin release under these two experimental conditions were taken into account. As judged from such differences expressed in absolute terms or relative to either the increment in the output attributable to the rise in D-glucose concentration from 5.6 to 16.7 mM or the close-to-basal value for insulin secretion found at 5.6 mM D-glucose, no obvious difference was found between (Ssun) rats and either

Figure 3. Difference in insulin output between islets exposed to either 10.0 mM 2-ketoisocaproate or 16.7 mM D-glucose and expressed, from left to right, in absolute terms and relative to either the increment in insulin release in response to a rise in D-glucose concentration from 5.6 to 16.7 mM or basal insulin output, all results being illustrated by reference to the mean value found in (Ssun) rats. The last set of columns on the right of the figure refers to the overall mean values derived from the former three comparisons. Same presentation as in Fig. 1.

Figure 4. Insulin content of islets first incubated for 90 min in the presence of D-glucose (G) 5.6, 8.3 or 16.7 mM, in the concomitant presence of 5.6 mM D-glucose (G) and 10.0 mM D-fructose (F), and in the presence of 10.0 mM 2-ketoisocaproate (KIC). Mean values (±SEM) refer to 74 separate determinations in all cases and are expressed relative to the average derived, in each individual experiment and in islets derived from the same group of rats, from the mean values recorded under the five experimental conditions prevailing during incubation. Also shown are the statistical significance (p) of differences between selected mean values.
As illustrated in Fig. 3, the results recorded in (Fsun) rats, however, averaged no more than 60.2±11.0% (df=84; p<0.004) of the mean corresponding values found in (Ssun) rats (100.0±7.6%; df=102). The results recorded in the former rats also differed significantly from those found in the (Fsaf) rats (94.5±9.3%; df=96; p<0.02) or (Fsal) rats (95.8±10.7%; df=81; p<0.025). Considering the lower absolute increments in insulin output provoked by a rise in D-glucose concentration from 5.6 to 16.7 mM in the islets from (Fsun) rats, as compared to (Ssun) rats (see above), the findings illustrated in Fig. 3 duly confirmed that there was no significant difference in the 4 groups of rats under consideration in terms of the islet secretory responsiveness to 2-ketoisocaproate.

Islet insulin content. As indicated in Table II, the mean insulin and protein content of pancreatic islets examined immediately after the isolation failed to differ significantly in the 4 groups of rats, with overall respective mean values of 450±34 µU/islet and 3.09±0.42 µg/islet (n=10 in both cases).

Likewise, in islets examined after incubation for 90 min at 37°C, the insulin content, expressed relative to the average of the mean islet insulin content found within each experiment and in the four groups of rats after incubation under the same experimental conditions, amounted to 93.6±2.6% (n=100) in the (Ssun) rats, 102.3±2.4% (n=85) in the (Fsal) rats, 103.4±3.0% (n=100) in the (Fsaf) rats, and 99.3±2.7% (n=85) in the (Fsaf) rats. Only the first of these four percentages differed significantly (p<0.002) from that recorded either in the (Fsun) rats or (Fsal) rats. Even so, the quite modest relative magnitude of such differences can obviously not be blamed to any major extent for differences in insulin output in the four groups of rats. Incidentally, the reference values used in these comparisons averaged 528±36 µU/islet (n=15).

As a rule, the mean values for islet insulin content recorded in each of the four groups of rats for islets first incubated under the same experimental conditions, when expressed relative to the average of the five mean values found in each experiment in islets from the same group of rats first incubated under five distinct experimental conditions (i.e. 528±43 µU/islet; n=12), failed to differ significantly from one another. As a matter of fact, among 30 comparisons, only two yielded a significant difference, such being the case in islets from (Ssun) and (Fsaf) rats first incubated at 5.6 mM D-glucose (df=35; p<0.05) and in islets from (Fsun) and (Fsal) rats first incubated at 16.7 mM D-glucose (df=35; p<0.02). In light of these findings, the overall mean values found in the four groups of rats were computed for each set of experimental conditions used during incubation. As illustrated in Fig. 4, the islet insulin content progressively decreased from 113.1±2.6% to 104.9±2.2% and 91.1±3.0% (n=74 in all cases) as the concentration of D-glucose during incubation increased from 5.6 to 8.3 and 16.7 mM. The insulin content was also lower (p<0.003) in islets first incubated at 5.6 mM D-glucose in the presence, as distinct from absence, of 10.0 mM D-fructose (102.6±2.0%; n=74). It was comparable in islets first exposed to either 16.7 mM D-glucose or 10.0 mM 2-ketoisocaproate (88.2±3.3%; n=74). These differences in islet insulin content obviously represented a mirror image of the differences in insulin output recorded during incubation under the same five distinct experimental conditions.

Table II. Insulin and protein content of freshly isolated islets.

<table>
<thead>
<tr>
<th>Rats</th>
<th>Protein content (µg/islet)</th>
<th>Insulin content (µU/islet)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ssun</td>
<td>3.6±0.91 (4)</td>
<td>444±69 (3)</td>
</tr>
<tr>
<td>Fsun</td>
<td>3.13±0.97 (2)</td>
<td>501±7 (2)</td>
</tr>
<tr>
<td>Fsaf</td>
<td>2.41±0.54 (3)</td>
<td>380±70 (3)</td>
</tr>
<tr>
<td>Fsal</td>
<td>3.00 (1)</td>
<td>514±99 (2)</td>
</tr>
</tbody>
</table>

Discussion

The mean close-to-basal value for insulin release recorded in islets incubated in the presence of 5.6 mM D-glucose was higher in the (Ssun) rats than in the other groups of rats. In the four groups of rats considered in this study, the mean values for such an insulin output were inversely related (r=−0.9825; n=4) to the mean corresponding values for the apparent caloric efficiency of the diet. The highest and lowest mean values for insulin release from islets exposed to 5.6 mM D-glucose also coincided, respectively, with the close-to-lowest and highest mean values recorded in the comparison between the increments in insulin output at 8.3/16.7 mM D-glucose (Fig. 2). These findings are compatible with the view that the apparent caloric efficiency of the diet participates in the regulation of the islet content in endogenous nutrients and, hence, in the relative magnitude of their secretory responsiveness to exogenous nutrients.

The comparison between the increments in insulin output caused by a rise in D-glucose concentration from 5.6 to 16.7 mM or by the addition of 10.0 mM D-fructose to a medium containing 5.6 mM D-glucose convincingly illustrated that the incorporation of large amounts of D-fructose in the diet failed to favour the secretory responsiveness to the ketohexose relative to that to the aldohexose (Fig. 1).

The dietary substitution of starch by D-fructose apparently lowered the postulated Km for the insulin secretory response to D-glucose (Fig. 2), an effect counteracted by the enrichment of the diet in long-chain polyunsaturated fatty acids, especially C20:5ω3 and C22:6ω3.

Lastly, as a rule, the dietary manipulations exerted little effects upon the insulinotropic efficiency of 2-ketoisocaproate, compared to that of D-glucose. At the most, the results illustrated in Fig. 3 suggest that the dietary substitution of starch by D-fructose decreased the insulinotropic efficiency of D-glucose, compared to that of 2-ketoisocaproate. The latter effect was again counteracted by the enrichment of the diet in either C18:2ω6 or C20:5ω3 and C22:6ω3.

It should be stressed that the changes in B-cell secretory behaviour encountered in the present study could not be blamed on any sizeable alteration in the insulin content of the islets. Moreover, the close parallelism observed between the amount of insulin released by the islets during incubation in the presence of different concentrations or combinations of nutrient secretagogues and the corresponding resulting changes in the insulin content of the islets further document the internal consistency of the present secretory data.

Taken as a whole, therefore, the present findings draw attention to three changes in the secretory behaviour of insulin-
producing cells caused, in sunflower oil-fed rats, by the dietary substitution of starch by D-fructose, namely a decrease in basal insulin output, an apparent lowering of the quoted Km for the insulinotropic action of D-glucose and an alteration in the insulinotropic efficiency of D-glucose relative to that of other nutrient secretagogues. Some of these secretory perturbations were opposed by the enrichment of the diet in long-chain polyunsaturated fatty acids, especially C20:5ω3 and C22:6ω3. To the extent that the latter dietary manipulation also opposed the insulin resistance caused in the present experimental design by the fructose-enriched diet, the above-mentioned changes in B-cell secretory behaviour could be attributable, in part at least, to such an insulin resistance considered as a major feature of the present animal model of metabolic syndrome.

Acknowledgments

We are grateful to C. Demesmaeker for secretarial help.

References