Abstract. The effects of chronic uremia and glucagon administration on glucagon-stimulable adenylcyclase in rat liver were assessed by determinations of adenyl cyclase activities, specific iodoglucagon binding, and the activity of the stimulatory regulatory component of adenyl cyclase. Glucagon-stimulated adenyl cyclase was reduced in uremia to 75–80% of control levels ($P < 0.05$), in the presence or absence of saturating levels of guanosine triphosphate (GTP) and $S^\prime$-guanylylimidodiphosphate [GMP-P(NH)P]. Although these changes were accompanied by a concomitant 20% reduction in sodium fluoride-stimulated activity, basal, GTP-, GMP-P(NH)P-, and manganese-dependent adenyl cyclase activities were unchanged. Using $[^{125}\text{I}}$-Tyr$^{10}$moniodoglucagon as a receptor probe, the number of high affinity glucagon-binding sites was reduced 28% ($P < 0.01$) in uremic as compared with control liver membranes. However, the affinity of these binding sites was unaltered. The S49 cyc$^-$reconstituting activity with respect to both GMP-P(NH)P- and isoproterenol plus GTP-stimulable adenyl cyclase was unchanged in membranes from uremic as compared with control rats. Intermittent glucagon (80–100 $\mu$g) injections administered at 8-h intervals to normal rats reproduced all of the above described effects of chronic experimental uremia on the adenyl cyclase system. It is concluded that changes in the hormone-stimulable adenyl cyclase complex in uremia and with glucagon treatment result primarily from a decrease in the number of hormone-specific receptor sites in hepatic plasma membranes. Since the changes in liver adenyl cyclase are qualitatively and quantitatively the same in glucagon-treated and uremic rats, it is suggested that these may be the result of the hyperglucagonemia of uremia. Further, the data reveal an unexpected dissociation between guanine nucleotide and sodium fluoride stimulation of adenyl cyclase. Possible causes for this dissociation based on the known subunit composition of cyclase coupling proteins are discussed.

Introduction

Abnormal carbohydrate metabolism is a frequent concomitant of chronic renal disease (1–3). As the result, in part, of the diminished renal clearance of glucagon, hyperglucagonemia is frequently observed in renal insufficiency (4, 5). These increased immunoreactive glucagon levels result from increases in a number of molecular weight species, each of which may have glucagon-like immunoreactivity if not biological activity (6–8). Glucagon-stimulable adenyl cyclase activity in liver of uremic rats has been found to be increased (9, 10). Other more indirect assessments have inferred a decreased activity of this enzyme in chronic uremia (11). In light of these discordant data and to determine the impact of the uremic state on the hormone-stimulable adenyl cyclase complex, we have used a monoiodinated glucagon as a probe of glucagon receptor density and affinity in membrane preparations of rat liver (12), and have now developed and validated reconstitution assays using S49 cyc$^-$lymphoma cell membranes to measure the activity of the stimulatory regulatory component which links the glucagon receptor to the catalytic component of adenyl cyclase. The effects of chronic uremia have been compared with the effects of intermittent glucagon administration in vivo on these parameters in rat liver. The implications of these findings for the regulation of hormone-stimulable adenyl cyclase activities are discussed.
Methods

Myokinase, creatine phosphokinase, creatine phosphate, ATP, cAMP, guanosine triphosphate (GTP), DNA, I, isoproterenol, and other biochemical reagents were obtained from Sigma Chemical Co. (St. Louis, MO). Cholic acid was also purchased from Sigma Chemical Co. and recrystallized from ethanol six times before use. [α-32P]ATP (25 Ci/mmol) was either purchased from the International Chemical and Nuclear Corp. (Irving, CA) or supplied by the Molecular Endocrinology Core Laboratory of the Baylor Center for Diabetes and Endocrinology Research. [3H]cAMP (~21 Ci/mmol sp act) was purchased from Amersham Corp. (Arlington Heights, IL). 5′ Guanylylimidodiphosphate [GMP-P(NH)P] was purchased from Boehringer Mannheim Biochemicals (Indianapolis, IN). DEAE-Sephadex was purchased from Pharmacia Fine Chemicals (Piscataway, NJ). Glucagon injected into animals was purchased from Sigma Chemical Co. or Eli Lilly & Co. (Indianapolis, IN). Glucagon used in adenyl cyclase assays and for iodination was a generous gift of Dr. W. W. Broomer (Eli Lilly & Co.). [125I]Tyro-2-monooiodoglucagon, prepared by iodination of native glucagon with carrier-free sodium [125I]iodide in the presence of iodogen followed by high pressure liquid chromatography purification using a μBondapak C18 column described by Rojas et al. (12), was supplied by the Molecular Core Laboratory of the Baylor Diabetes and Endocrinology Research Center.

Female Sprague-Dawley rats (150–200 g) from TIMCO (Houston, TX) were used in these studies. Animals were made uremic by unilaterial nephrectomy and ligation of two branches of the contralateral renal artery as described previously (13, 14). The animals were maintained on a high-protein diet and then sacrificed by decapitation after 7 d. Blood-urea nitrogen (as milligrams per 100 milliliters whole blood) was 57±4 (mean±SE) in the uremic group and 14±2 in the control group.

Glucagon-treated animals were prepared as described in Results.

Membrane preparations

The livers from uremic, normal, or glucagon-treated rats were quickly removed, placed on ice, and the membranes prepared according to the procedure of Neville (15) as modified by Pohl et al. (16). The membranes were then stored at ~80°C. Cyclic-murine lymphoma S49 cells were grown and membranes (cyc membranes) were prepared by Dr. Ravi Iyengar (Department of Cell Biology, Baylor College of Medicine) using procedures described elsewhere (17, 18).

Adenylyl cyclase assays

Liver membranes (8–12 µg protein) were incubated for 10 min at 30°C in a final volume of 50 µl containing 0.1 mM [α-32P]ATP (1,000–1,500 cpm/pmol), 5 mM MgCl2, 1 mM EDTA, 1 mM [3H]cAMP (~10,000 cpm), 25 mM Tris-HCl, the indicated additives, and an ATP-regenerating system consisting of 0.02 mg/ml myokinase, 0.2 mg/ml creatine phosphokinase, and 20 mM creatine phosphate. The incubations were terminated by the addition of 100 µl of a stop solution consisting of 40 mM ATP, 10 mM cAMP, and 1% sodium dodecyl sulfate. [32P]cAMP formed was quantitated by the method of Salomon et al. (19) as modified by Bockaert et al. (20). Both the ATP and the reagents used to regenerate it during the assays were purified from contaminating “GTP-like” materials as described elsewhere (21, 22).

1. Abbreviations used in this paper: GMP-P(NH)P, 5′ guanylylimidodiphosphate; GTP, guanosine triphosphate.

Glucagon-binding assays

Membranes (2–5 µg protein) were incubated for 20 min at 32.5°C in a final volume of 100 µl of a medium containing varying concentrations of [125I]monooiodoglucagon (0.20–3.0 nM), 25 mM Tris-HCl, pH 7.5, 1.0 mM EDTA, 0.1% bovine serum albumin (BSA), and other additions as indicated. At the end of the incubation, the reaction mixtures were diluted with 5 ml of ice-cold 25 mM Tris-HCl, pH 7.5, and 0.1% BSA (mucilagum), and filtered rapidly through 0.45-µm cellulose acetate filters (Nalgene Oxoid, Columbia, MD) that had been presoaked overnight in 10% BSA. The tubes were rinsed once with 5 ml of ice-cold buffer and the rinse filtered through the Oxoid filters, which were then immediately washed with a final 5-ml aliquot of the ice-cold buffer. The filters were counted in a Searle Analytical gamma counter at a counting efficiency of 50%. To determine the nonspecific binding, membranes were incubated under identical conditions with [125I]iodoglucagon in the presence of 3.0 µM unlabeled glucagon. Specific binding was determined by subtracting nonspecific bound counts from the total bound counts. The data obtained were analyzed according to Scatchard (23) to determine both the dissociation constant (Kd) and the number of specific binding sites on the membranes tested.

Measurement of regulatory component activity of liver membranes by cholate extraction and reconstitution into cyc membranes

Extraction of regulatory component activity and selective inactivation of coextracted catalytically active adenyl cyclase activity. Membranes were mixed with extraction medium to give a final concentration of 1% cholate, 25 mM Na-HEPES, pH 8.0, 1.0 mM EDTA, and 10 mM MgCl2. Membrane concentration during extraction was 5–7.5 mg/ml. The membrane-cholate mixtures thus obtained were kept on ice for 60 min with occasional stirring. The EDTA concentration was increased to 10 mM to cholate excess MgCl2 and the mixtures centrifuged in the cold at 100,000 g for 60 min. The supernatants were then warmed at 30°C for 5 min yielding the cholate extracts in which regulatory component activity was measured by reconstitution of GMP-P(NH)P or isoproterenol plus GTP-stimulated adenyl cyclase activities in cyc membranes. As shown under Results, the above procedure results in a recovery of ~75% of the total regulatory component activity measurable in the membrane-cholate mixtures.

Reconstitution of GMP-P(NH)P-stimulated activity in cyc membranes. Cholate extracts were diluted 25-fold or more with dilution medium consisting of 1.5 M KCl and 40 mM β-mecaptoethanol. Cyc-reconstituting activity was then assayed by adding 10 µl of the diluted, warmed cholate extracts to 10 µl of cyc-membrane suspension (25–35 µg in 1 ml diethanol and 25 mM Na-HEPES, pH 8.0), keeping this mixture in the cold for 10–30 min, and assaying for GMP-P(NH)P-stimulated cyc adenyl cyclase activity by further addition of 30 µl of assay reagents to give, in a final volume of 50 µl, 0.1 mM [α-32P]ATP (specific activity 2,000 cpm/pmol), 10 mM MgCl2, 1.0 mM [3H]AMP (~10,000 cpm/assay), 1.0 mM EDTA, the ATP-regenerating system described above for adenyl cyclase assays, 25 mM Tris-HCl, and 100 µM GMP-P(NH)P. The mixtures were incubated for 40 min at 30°C, stopped, and analyzed for [32P]cAMP formed as described (19).

Reconstitution of cyc isoproterenol plus GTP-stimulated activity. Cholate extracts were diluted as above at least fivefold in dilution medium. Aliquots (20 µl) were then incubated for 10 min at 30°C with cyc membranes (250–350 µg protein) in a final volume of 100 µl containing 0.5 mM ATP, 2.0 mM MgCl2, 1 mM EDTA, the ATP-regenerating system described above, and 0.1 mM GTP. This phase of the
procedure is the reconstitution proper. 10-µl aliquots of the reconstituted mixtures were then transferred to tubes having 30 µl of a mixture containing 16.7 mM MgCl₂, 1.67 mM EDTA, 1.67 mM [³H]cAMP (~10,000 cpm), the above described ATP-regenerating system, 41.7 mM Tris-HCl, pH 7.5, and 1.67 µM 1-isoproterenol. After 30 min at 30°C, the reconstituted isoproterenol plus GTP-stimulated activity was monitored by postaddition of 10 × 10⁶ cpm of high specific activity [α-³²P]ATP contained in 10 µl. The reactions (final concentration of ATP was 0.1 mM and of MgCl₂ was 10 mM) were stopped after 10 min and the [³H]cAMP formed was quantitated as described.

By using these assay conditions and at the dilutions of cholate extract employed, the cyc⁻ reconstituted activities were proportional to the microgram quantities of cholate extract protein added. The results of these reconstitution assays are expressed as cyc⁻-reconstituted activities in picomoles of cAMP formed per 40-min assay in the case of reconstitution of GMP-P(NH)P-stimulated activities, or as picomoles of cAMP formed per 10-min assay in the case of reconstitution of isoproterenol plus GTP-stimulated activities. In some instances, these activities were corrected for by dividing the above result by the milligram quantity of cholate extract protein added, and thereby yielding specific activities of cyc⁻-reconstituting activities which are expressed as nanomoles of cAMP formed per 40- or 10-min assays per milligram of cholate extract protein.

Variability of results
The activity values obtained for a given membrane cholate extract were reproducible from day to day within 10% limits. Replicates within a given assay agreed within 5%. As prepared, cholate extracts could be stored at −80°C without loss of activity for up to 4 mo, the longest time tested.

Although cholate extracts from different control liver membrane batches gave very similar cyc⁻-reconstituting activities (as tested in a single assay), the absolute values of cyc⁻-reconstituted activities obtained (i.e., picomoles of cAMP formed under the assay conditions) varied by as much as fivefold depending on the batch of cyc⁻ membranes used. This is owing to the fact that reconstituted activities depend on the quality of the cyc⁻ adenyl cyclase system used to assay for regulatory component. In spite of these variations which relate to variability in the growth rate of cells and the cell density at the time of harvest, reconstituted activities were always proportional to micrograms of cholate extract protein added. Thus, the cyc⁻ membranes used provided linear assays of reconstituting activity. For these reasons, all studies in which regulatory component activities from various liver membrane preparations are compared were measured in single experiments using a single batch of cyc⁻ membranes. Protein was determined by the method of Lowry et al. (24).

Results
Effects of uremia. The impact of chronic uremia on glucagon-stimulable adenyl cyclase in liver was assessed in membrane preparations of rats made chronically uremic by surgical techniques described previously (13, 14). Although basal adenyl cyclase was not different in uremic as compared with control membranes (9.9±0.9, 10.3±2.8 pmol cAMP formed/min per mg membrane protein, respectively; ±SD, n = 6), glucagon-stimulable activity was decreased 27% in uremia (53.2±4.5, 39.1±6.1; P < 0.05) (Table I). No significant effect on the concentration of glucagon required for half-maximal stimulation

<table>
<thead>
<tr>
<th>Additions to assays</th>
<th>Control</th>
<th>Uremic</th>
</tr>
</thead>
<tbody>
<tr>
<td>Glucagon + GTP</td>
<td>249.4±15.0</td>
<td>207.8±5.4*</td>
</tr>
<tr>
<td>NaF</td>
<td>226.5±27.6</td>
<td>183.8±11.8*</td>
</tr>
<tr>
<td>MnCl₂</td>
<td>36.4±3.2</td>
<td>29.9±4.3</td>
</tr>
</tbody>
</table>

Membranes were incubated and adenyl cyclase activity determined as described under Methods in the absence and presence of 10 µM GTP, 10 µM GMP-P(NH)P, 10 mM NaF, 10 mM MnCl₂, and 1 µM glucagon separately or in combination as indicated. If MnCl₂ was added, MgCl₂ was omitted. Membranes were incubated for 10 min at 32.5°C as described more completely in Methods. Values shown are the mean±SD of at least six experiments. * P < 0.05

was noted in uremic as compared with control rats (Fig. 1). The stimulations by saturating glucagon in the presence of GTP and GMP-P(NH)P were reduced by 14 and 17%, respectively (P < 0.05) (Table I). Although the NaF-stimulated activity was reduced 19% (P < 0.05), GTP, GMP-P(NH)P, and MnCl₂ ac-

![Figure 1. Concentration-effect curves for glucagon stimulation of adenyl cyclase activity in liver membranes from control (---) and uremic (---) rats. Activities were assayed for 10 min at the indicated concentrations of glucagon using, per assay, 5.2 and 7.3 μg of liver membrane protein from control and uremic rats, respectively. For the rest of the conditions, see Methods.](image-url)
tivities were not different in uremic as compared with control preparations.

The activity of the guanine nucleotide-binding α-subunit of regulatory component was next examined in control and uremic rats by using two methods of assessment. First, a reconstitution of GMP-P(NH)P-stimulable adenylyl cyclase activity in membranes of S49 cYC- lymphoma cell membranes was performed by using regulatory component extracted from rat liver membranes with cholate. Second, the ability of added regulatory component extracted from liver membranes to confer, in the presence of GTP, isoproterenol stimulation to adenylyl cyclase in S49 cYC- cell membranes was assessed. The optimal conditions required for these reconstitution assays were investigated and validated. These studies indicated that reconstitution of GMP-P(NH)P-stimulated cYC adenylly cyclase activity was best observed in the presence of 20 mM β-mercaptoethanol and 0.75 M KCl, which both reduced the lag in appearance in reconstituted activity and led to a constant accumulation of cAMP for 60–80 min. Assays were performed routinely for 40 min. Optimal extraction of reconstituting activity from hepatic membranes was obtained in the presence of 1% cholate, 1 mM EDTA, and 10 mM MgCl_2 followed by chelation of excess MgCl_2 by postaddition of 11 mM EDTA. Coextracted cyclizing activity was inactivated by heating without major loss (10%) of the reconstituting activity. Optimal reconstitution of isoproterenol plus GTP-stimulated activities required cholate concentrations to be <0.1% during the first reconstitution phase of assay. The reconstituted adenylyl cyclase activity was augmented by KCl addition in the presence of low but not high cholate concentrations. The dependency on time of the reconstitution of isoproterenol-stimulable adenylyl cyclase activity and S49 cYC- cell membranes was such that hormonal reconstitution incubations were carried out for 10 min rather than 40 min. Under these conditions, the reconstitution of the hormonally sensitive adenylyl cyclase activity was linearly dependent on the quantity of regulatory component (cholate extract added) provided the quantity of liver membrane-derived protein in the cholate extract added to the reconstitution assays was <1 µg.

We investigated the efficiency with which regulatory component was extracted from liver membranes using the procedures described above. Nielsen et al. (25) have shown that, in cYC- cell membranes, reconstitution of GMP-P(NH)P-stimulated activation of adenylyl cyclase occurs not only with fully solubilized regulatory component, but also with partially solubilized component such as would be present in an adsorbed form on membranes treated with 1% cholate. We, therefore, measured "total reconstituting activity" by mixing the membrane-cholate mixtures with cYC- membranes and assessing the resulting GMP-P(NH)P-stimulated activity, which reflects totally reconstituted cYC activity, undisturbed liver membrane activity, and some as yet undefined level of liver membrane-bound regulatory component on cYC- membrane-bound adenylyl cyclase. We found that before warming, cholate extracts contained ~75% of the "maximally" measurable regulatory component activity (not shown). Based on these preliminary studies, it is apparent that (a) the bulk of the regulatory component activity can be extracted from liver membranes, (b) the extracts obtained can be warmed to inactivate endogenous catalytic activity without substantial loss of regulatory component activity, and (c) reconstituted activities can be used to quantitate regulatory component activity, provided that the cholate is diluted sufficiently and that both KCl and β-mercaptoethanol are added to accelerate the reconstitution process.

Using these techniques, regulatory component activity was determined in liver membrane preparations of uremic and control rats. Both the "extractability" and cYC- reconstituting activity of regulatory component activity, as assessed by reconstitution of GMP-P(NH)P-stimulable adenylyl cyclase activity, were not different in membranes of uremic as compared with control rats (Table II). Similarly, the capacity of regulatory component from membranes of uremic rats to reconstitute cYC isoproterenol plus GTP-stimulated activity was also unchanged when compared with controls. Thus, using these two assessments of regulatory component activity, the activity of this coupling protein in uremic membranes was found to be unaltered as compared with membranes from control animals.

Because the foregoing data suggest potential alterations of

<p>| Table II. Regulatory Component Activities in Liver Membranes from Control and Uremic Rats |</p>
<table>
<thead>
<tr>
<th>Membranes</th>
<th>GMP-P(NH)P</th>
<th>Iso plus GTP</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>cAMP formed/40 min/ mg cholate extract protein</td>
<td>cAMP formed/10 min/ mg cholate extract protein</td>
</tr>
<tr>
<td>Control</td>
<td>13.9±0.3 (n = 6)</td>
<td>1.03±0.11 (n = 6)</td>
</tr>
<tr>
<td>Uremic</td>
<td>14.5±0.8 (n = 6)</td>
<td>1.08±0.15 (n = 6)</td>
</tr>
</tbody>
</table>

Liver membranes from control and uremic rats (5–7 mg protein/ml) were extracted as described under Methods, yielding cholate extracts of 2.5–3.1 mg protein/ml. Each of the extracts was then assayed in duplicate at three different dilutions with cYC membranes for reconstitution of GMP-P(NH)P-stimulable and isoproterenol plus GTP-stimulated activities. Final dilutions of cholate extracts during the reconstitution phases of incubations were 250, 500, and 1000 when GMP-P(NH)P-stimulated activities were reconstituted and 50, 100, and 200 when isoproterenol plus GTP activities were reconstituted. This corresponded to input cholate extract proteins of 0.5–0.62 µg/32.5 µg cYC- membrane protein at the lowest dilution tested and assured proportionality in the assay. Specific activities for reconstituting cYC adenylly cyclase activities were then calculated, confirmed to be independent of sample dilution under the assay conditions used, and averaged to give the values reported in the table. For the rest of the conditions, see Methods.

* In all final adenylly cyclase assay stages of incubations, cYC- membrane protein was 32.5 µg. Cyclic AMP due to cYC- membranes alone was 0.1 pmol/40 min per assay.

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glucagon receptor function in liver membranes of uremic rats, the number of glucagon receptors and their affinity were examined using monoiodinated glucagon. Binding studies with this highly purified $[^{125}]$iodoglucagon were performed in hepatic membrane preparations from control and uremic rats, and the data were analyzed by the method of Scatchard (23). The data obtained (Fig. 2) indicate the existence in liver membranes of an apparently single population of high affinity binding sites for the probe. No differences in receptor affinity for this radio-labeled ligand could be detected in membranes of uremic as compared with control animals (Table III). However, the quantity of specific $[^{125}]$iodoglucagon-binding sites was reduced from 5.77±0.18 to 4.13±0.18 pmol of glucagon bound/mg membrane protein ($P < 0.001 ; n = 6$).

**Effect of glucagon administration.** It has been suggested that the hyperglucagonemia of chronic renal failure may alter glucagon-responsive adenylyl cyclase activity (9–11). For this reason, potential effects of intermittent glucagon injections on glucagon-sensitive adenylyl cyclase in rat liver were investigated. Liver membranes were prepared from rats administered 80–100 μg glucagon (subcutaneous) at 8-h intervals for 3 d; the properties of their adenylyl cyclase system were compared with those in control liver membranes prepared in parallel. As shown in Tables IV–VI and Figs. 3 and 4, glucagon administration reproduced both qualitatively and quantitatively the results obtained in liver membranes of uremic rats: glucagon and sodium fluoride ac-

### Table III. Glucagon Receptor Density and Affinity in Liver Membranes of Control and Uremic Rats

<table>
<thead>
<tr>
<th>Animal state</th>
<th>$K_D$</th>
<th>$B_{max}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Control</td>
<td>1.85 (1.26–3.46)</td>
<td>5.77±0.18 pmol/mg</td>
</tr>
<tr>
<td>Uremia</td>
<td>1.87 (1.31–3.27)</td>
<td>4.13±0.18* pmol/mg</td>
</tr>
</tbody>
</table>

Liver membranes from control and uremic rats were prepared and incubated as described in Methods. The number and affinity of specific $[^{125}]$-Tyr$^9$monoiodoglucagon-binding sites were calculated by means of Scatchard analysis of saturation curves of purified monoiodoglucagon, as shown in Fig. 2. Receptor density values ($B_{max}$) are the mean±SE ($n = 6$); 95% confidence limits for the $K_D$ values are given in parentheses.

* $P < 0.001$ compared with control.

activities of adenylyl cyclase were reduced, but basal, GTP-, GMP-P(NH)$_2$-, and manganese-dependent activities were unchanged; the α-subunit activity of the stimulatory regulatory component was unaltered; and glucagon receptor levels but not affinity were reduced.

### Table IV. Adenylyl Cyclase Activities of Liver Membranes from Control and Glucagon-treated Rats

<table>
<thead>
<tr>
<th>Additions to assays</th>
<th>Control</th>
<th>Glucagon-treated</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>pmol cAMP/min per mg</td>
<td>pmol cAMP/min per mg</td>
</tr>
<tr>
<td>None</td>
<td>8.1±1.4</td>
<td>7.5±0.4</td>
</tr>
<tr>
<td>GTP</td>
<td>25.5±2.0</td>
<td>28.8±4.8</td>
</tr>
<tr>
<td>GMP-P(NH)$_2$</td>
<td>74.7±5.6</td>
<td>72.4±8.2</td>
</tr>
<tr>
<td>Glucagon</td>
<td>59.3±2.9</td>
<td>46.2±3.5*</td>
</tr>
<tr>
<td>Glucagon + GTP</td>
<td>148.8±7.5</td>
<td>119.4±1.8*</td>
</tr>
<tr>
<td>Glucagon + GMP-P(NH)$_2$</td>
<td>200.5±9.6</td>
<td>159.6±3.8*</td>
</tr>
<tr>
<td>NaF</td>
<td>181.8±11.6</td>
<td>147.4±4.5*</td>
</tr>
<tr>
<td>MnCl$_2$</td>
<td>29.4±1.3</td>
<td>28.9±1.6</td>
</tr>
</tbody>
</table>

Adenylyl cyclase activity was determined in membranes obtained from control rats and rats injected with 80 μg glucagon subcutaneously every 8 h for 3 d and sacrificed 3 h after the final injection. As described in Methods, membranes were incubated for 10 min at 32.5°C in the absence and presence of 1 μM glucagon, 10 μM GTP, 10 μM GMP-P(NH)$_2$, 10 mM NaF, or 10 mM MnCl$_2$ separately or in combination as indicated. If MnCl$_2$ was added, MnCl$_2$ was omitted. Values shown are the mean±SD of at least three experiments.

* $P < 0.05$. 

![Figure 2. Scatchard analysis of $[^{125}]$iodoglucagon binding to liver membranes of control and uremic animals. $[^{125}]$iodoglucagon binding was determined as described under Methods. The values of free and specifically bound $[^{125}]$iodoglucagon obtained were plotted according to Scatchard (23). The inset shows the saturation curves from which the Scatchard lines for normal (——) and uremic (——) membranes were derived. Each point on the saturation curve represents the mean of three determinations. B/F, bound/free.](image-url)
Table V. Regulatory Component Activities in Liver Membranes from Control and Glucagon-treated Rats

<table>
<thead>
<tr>
<th>Membranes</th>
<th>GMP-P(NH)P</th>
<th>Iso plus GTP</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>nmol cAMP formed/40 min</td>
<td>nmol cAMP formed/10 min</td>
</tr>
<tr>
<td>Control</td>
<td>10.4±0.9 (n = 12)</td>
<td>0.65±0.04 (n = 9)</td>
</tr>
<tr>
<td>Glucagon-treated‡</td>
<td>10.5±0.4 (n = 6)</td>
<td>0.66±0.05 (n = 6)</td>
</tr>
</tbody>
</table>

Liver membranes from control and glucagon-treated rats (5–7 mg protein/ml) were extracted as described under Methods yielding cholate extracts of 2.6–2.9 mg protein/ml. Each of the extracts was then assayed in triplicate (controls) or duplicate (glucagon-treated) at three different dilutions for reconstitution of GMP-P(NH)P-stimulated and isoproterenol plus GTP-stimulated c yc- adenylyl cyclase activities. Specific activities for reconstitution of c yc- adenylyl cyclase were calculated, confirmed to be independent of sample dilution, and averaged to give the values reported in the table. For further details, see Methods.

* In all final assay stages of incubations, c yc- membrane protein was 27.8 µg and cAMP formed due to c yc- membranes alone was subtracted from conversions obtained in the presence of cholate extracts prior to calculation of specific reconstituting activities.

‡ Animals were treated with 80 µg glucagon subcutaneously every 8 h for 3 d and killed 3 h after the last injection.

Discussion

Altered glucagon stimulation of liver adenylyl cyclase in uremia. Disparate conclusions regarding hepatic glucagon-sensitive adenylyl cyclase activity in chronic uremia have been reached (9–

Table VI. Glucagon Receptor Density and Affinity in Liver Membrane of Control and Glucagon-injected Rats

<table>
<thead>
<tr>
<th>Animal state</th>
<th>K_D</th>
<th>B_max</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>nM</td>
<td>pmol/mg</td>
</tr>
<tr>
<td>Control</td>
<td>1.83</td>
<td>5.77±0.18</td>
</tr>
<tr>
<td></td>
<td>(1.45–2.59)</td>
<td></td>
</tr>
<tr>
<td>Glucagon-injected</td>
<td>2.10</td>
<td>4.04±0.12*</td>
</tr>
<tr>
<td></td>
<td>(1.78–2.62)</td>
<td></td>
</tr>
</tbody>
</table>

Liver membranes were prepared from control animals and animals injected with 80 µg glucagon thrice daily for 3 d. The membranes were incubated as described in Methods and Scatchard analysis was performed for both types of membranes on saturation curves of [125I]iodoglucagon. Calculated receptor affinities and densities are given below. B_max values are the means±SE for at least three separate experiments; 95% confidence limits provided for the K_D values are given in parentheses.

* P < 0.001 compared with control.
Although the basis for these discrepant findings is unclear, the results of the present study show clearly that the glucagon-stimulable adenylyl cyclase in liver of uremic rats is impaired in a relatively complex manner (Fig. 1, Table I). All forms of glucagon-stimulated activities, both without and with added guanine nucleotides, as well as fluoride-stimulated activity were reduced 20–30% in uremia. In contrast, basal, manganese-, and guanine nucleotide-stimulated activities per se were unaltered.

Glucagon-receptor number but not affinity was decreased in uremia (Fig. 2, Table III). Analysis of the α-subunit of the regulatory component activity, as assessed by reconstitution of the S49 cyc− adenylyl cyclase, showed no change. Since the reduction in glucagon-specific binding sites (~28%) correlated well with the reduction in glucagon-stimulated adenylyl cyclase (~27%), the data strongly suggest that the decrease in glucagon-stimulable adenylyl cyclase activity may be the result of a loss or inactivation of glucagon receptors without significant changes in the remainder of the enzyme complex.

There are three mechanisms which might potentially account for the loss of glucagon-responsive adenylyl cyclase in liver membranes from chronically uremic rats. First, a nonspecific or toxic effect of the uremic state might result in a depressed activity of the enzyme complex. This would appear to be somewhat unlikely since the effect is localized specifically to the number of glucagon-binding sites. Such a change would perturb alterations in receptor affinity, in coupling factors, or in the maximal catalytic capacity of the enzyme would be unexpected, but cannot be entirely excluded. Second, a heterologous desensitization of the glucagon-responsive adenylyl cyclase complex may have been produced as a result of the uremic state. In other tissues, alterations of regulatory component activity or of the kinetics of radioligand binding to the hormone receptor have been observed or suggested as the mechanism accounting for heterologous desensitization (26–28). Marked elevations of parathyroid hormone are a clear concomitant of the chronically uremic state (29–31). Similarly, we have reported elevated levels of catecholamines and a loss of responsiveness of the epinephrine-stimulable adenylyl cyclase in skeletal muscle of chronically uremic rats (32). It is therefore possible that primary alterations in either the epinephrine- or the parathyroid hormone-stimulable adenylyl cyclases or in both may account for the decreased glucagon responsiveness in uremia (33, 34). However, this possibility also appears to be somewhat unlikely since, as we show here, primary alterations both of regulatory component activity and in glucagon-receptor affinity are not observed. Finally, chronic uremia is associated with hyperglucagonemia (4–8), and changes in glucagon-stimulated cyclase activity might be the result primarily of the elevated circulating glucagon levels. We, therefore, tested whether or not glucagon administration to rats might mimic in any way the changes in adenylyl cyclase regulation observed in uremia. Glucagon-specific membrane receptors were decreased to a similar extent as in uremia; glucagon- and sodium fluoride-stimulated but not basal, GTP-, GMP-P(NH)P-, or manganese-dependent adenylyl cyclase activities were reduced to an extent similar to that observed in chronic uremia. There were no changes observed in α-subunit function of the regulatory component as suggested by reconstitution assays. Taken as a whole, these findings in glucagon-treated animals are entirely consistent with a glucagon-induced desensitization of the glucagon-stimulable adenylyl cyclase in rat liver, the decreased adenylyl cyclase activity in response to glucagon being the result primarily of the decreased glucagon-receptor density following glucagon administration (35–37). Since the data obtained from the glucagon-injected animals are in every way similar to the data obtained on the same hormone-stimulable adenylyl cyclase in liver of uremic rats, it would seem reasonable to conclude that the changes in liver of chronically uremic animals may derive primarily from a glucagon-induced desensitization of the hormone-sensitive adenylyl cyclase complex rather than from a nonspecific toxic effect of the uremic state.

Altered fluoride stimulation of adenylyl cyclase in uremia and glucagon treatment. Decreased sodium fluoride-stimulated adenylyl cyclase activity without concomitant changes in either basal, manganese-dependent, and GTP- or GMP-P(NH)P-stimulated activity was noted in hepatic membranes from both uremic and glucagon-treated as compared with control animals. Such a dissociation between nucleotide-stimulated as compared with fluoride-stimulated activities has not been reported previously. Both sodium fluoride and guanine nucleotides appear to exert their effects by interacting, as do hormone receptors, with the regulatory component (38–42). Detailed studies by Gilman and collaborators (43–46) have shown that the stimulatory regulatory component is formed of at least two distinct subunits: first, an α-subunit which is the substrate for ADP-ribosylation (47) and which regulates the catalytic activity of adenylyl cyclase per se, and second, a β-subunit for which no specific function has yet been assigned with certainty. The cyc− S49 cell membranes used lack only the α-subunit of the regulatory component (48, 49), but do contain the β-subunit. Since activation of regulatory component is dependent on or at least associated with its dissociation into the individual subunits (50, 51), the reconstitution assay used actually assesses the activated α-subunit fraction but not the β-subunit. Guanine nucleotides exert their action on adenylyl cyclase by interacting with and binding to the α-subunit of the regulatory component (50, 51). However, while sodium fluoride exerts its action on adenylyl cyclase at the level of the regulatory component (46), the specific subunit at which it interacts is not known, nor is the precise mechanism by which fluoride activates the regulatory component.

The results of the present study show clearly that both guanine nucleotide regulation of adenylyl cyclase, as assessed in the absence of hormone, and the cyc− reconstituting activity are unaltered in uremic and glucagon-treated as compared with control liver membranes. These findings strongly indicate that the α-subunit of the regulatory component is unaltered. Since sodium fluoride but not nucleotide regulatory activity was decreased in uremic as compared with control liver membranes,
it is possible that this reflects an alteration either of the level of another subunit or of the capacity of another subunit to interact with the α-subunit of the stimulatory regulatory component. This other subunit may be the β-subunit, or it may well be an as yet undefined factor regulating sodium fluoride but not guanine nucleotide stimulation of cyclic AMP formation. It is of interest that treatment of MDCK cells with glucagon results in a loss of glucagon- and sodium fluoride-stimulated but not basal or guanine nucleotide-regulated adenyl cyclase activities or in cyc-
reconstituting activity (52). These findings are essentially identical to the pattern of alterations in adenyl cyclase activities reported here. Thus, the dissociation between guanine nucleotide and sodium fluoride-stimulated activity observed in this study may not be a phenomenon restricted solely to uremia or to rat liver and may therefore reflect a more common phenomenon of adenyl cyclase regulation.

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