Interaction of Guanosine Cyclic 3',5'-Phosphate Dependent Protein Kinase with lin-Benzoadenine Nucleotides†

Deepak Bhatnagar,† David B. Glass,‡ Robert Roskoski, Jr.,*† Ralph A. Lessor,† and Nelson J. Leonard∥

Department of Biochemistry, Louisiana State University Medical Center, New Orleans, Louisiana 70119. Department of Pharmacology, Emory University School of Medicine, Atlanta, Georgia 30322, and Departments of Chemistry and Biochemistry, University of Illinois, Urbana, Illinois 61801

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ABSTRACT: Using the activated cGMP-dependent protein kinase in the presence of the phosphorylatable peptide [[Ala34]histone H2B-(29-35)], we found that lin-benzoadenosine 5' -diphosphate (lin-benzo-ADP) was a competitive inhibitor of the enzyme with respect to ATP with a $K_i$ (22 $\mu$M) similar to the $K_d$ (20 $\mu$M) determined by fluorescence polarization titrations. The $K_i$ for lin-benzo-ADP determined in the absence of the phosphorylatable peptide, however, was only 12 $\mu$M. ADP bound with lower affinity ($K_i = 169$ $\mu$M; $K_d = 114$ $\mu$M). With [Ala34]histone H2B-(29-35) as phosphoryl acceptor, the $K_m$ for lin-benzo-ADP was 29 $\mu$M, and that for ATP was 32 $\mu$M. The $V_{max}$ with lin-benzo-ATP, however, was only 0.06% of that with ATP as substrate. Binding of lin-benzo-ADP to the kinase was dependent upon a divalent cation. Fluorescence polarization revealed that Mg$^{2+}$, Mn$^{2+}$, Ca$^{2+}$, Ni$^{2+}$, Sr$^{2+}$, and Ba$^{2+}$ supported nucleotide binding to the enzyme; Ca$^{2+}$, Sr$^{2+}$, and Ba$^{2+}$, however, did not support any measurable phosphotransferase activity. The rank order of metal ion effectiveness in mediating phosphotransferase activity was Mg$^{2+}$ > Ni$^{2+}$ > Co$^{2+}$ > Mn$^{2+}$. Although these results were similar to those observed with the cAMP-dependent protein kinase [Hartl, F. T., Roskoski, R., Jr., Rosendahl, M. S., & Leonard, N. J. (1983) Biochemistry 22, 2347], major differences in the $V_{max}$ with lin-benzo-ATP as substrate and the effect of peptide substrates on nucleotide (both lin-benzo-ADP and ADP) binding were observed.

The two cyclic nucleotide dependent protein kinases, namely, the adenosine cyclic 3',5'-phosphate (cAMP) dependent and the cGMP-dependent protein kinases, have been purified to homogeneity, and many characteristics have been defined. A comparison of the subunit structure, physicochemical properties, kinetic characteristics, and substrate specificity of these two protein kinases has been made (Gill et al., 1977; Lincoln et al., 1977; Takai et al., 1976; Foster et al., 1981; de Jonge et al., 1983; Gill, 1977, 1978, 1983) and are supported by recent sequence data (Hashimoto et al., 1981, 1982; Takio et al., 1982; Titani et al., 1983). These determinations have used the fluorescent lin-benzo-ATP as substrate and the effect of peptide substrates on nucleotide (both lin-benzo-ADP and ADP) binding were observed.

(C) subunits (Corbin et al., 1978) as follows:

$$R_5C_5 + 4cAMP \rightarrow R_5(cAMP)_4 + 2C$$

whereas cGMP-dependent protein kinase is a dimer that is activated without subunit dissociation (Corbin & Doskeland, 1983; Gill et al., 1977; Lincoln et al., 1977; Takai et al., 1976):

$$cGMPdPK, + 4cGMP \rightarrow (cGMPdPK)_2(cGMP)_4$$

Walter et al. (1980) have reported that antisera against the cGMP-dependent protein kinase does not cross-react with either catalytic or regulatory subunits of the type I or type II cAMP-dependent protein kinases and vice versa. Although amino acid sequence data indicate that a few regions of the two kinases do not correspond to each other, the ATP binding sites, the cyclic nucleotide binding sites, and the sites of autophosphorylation in the hinge regions show from 25 to 45% sequence identity between the two enzymes (Hashimoto et al., 1981, 1982; Takio et al., 1982; Titani et al., 1983). These results suggest that similarities exist in the functionally important regions such as the ATP-binding site. In order to ascertain the degree of these hypothesized similarities between the cAMP-dependent and cGMP-dependent protein kinases, we have used steady-state kinetics and a fluorescence polarization titration technique for characterizing the interaction of nucleotides with the cGMP-dependent protein kinase. For these determinations we have used the fluorescent lin-benzo-adenosine nucleotides, "stretched-out" analogues of adenosine.

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∥Louisiana State University Medical Center.
*Emory University School of Medicine.
†University of Illinois.

1Abbreviations: cGMP, guanosine cyclic 3',5'-phosphate; cAMP, adenosine cyclic 3',5'-phosphate; lin-benzo-ADP, lin-benzoadenosine 5'-diphosphate; Ser-peptide, Leu-Arg-Ala-Ser-Leu-Gly; [Ala34]-histone H2B-(29-35), Arg-Lys-Arg-Ser-Arg-Ala-Glu; histone H2B-(29-35), Arg-Lys-Arg-Ser-Arg-Lys-Glu; Mops, 3-(N-morpholino)propanesulfonic acid; MES, 2-(N-morpholino)ethanesulfonic acid; Tris, tris(hydroxymethyl)aminomethane.
nucleotides (Leonard et al., 1978). We have previously shown that lin-benzo-ADP is a competitive inhibitor of the catalytic subunit of the cAMP-dependent protein kinase with respect to ATP with a $K_i$ (8.0 $\mu$M) similar to the $K_i$ for ADP (9.0 $\mu$M) (Hartl et al., 1983). The fluorescence polarization studies also showed that the catalytic subunit of that enzyme and the type II holoenzyme bind lin-benzo-ADP rigidly; lin-benzo-ATP, moreover, is a very good substrate for the phosphotransferase activity of the protein kinase with peptides, water, or type II regulatory subunit as phosphoryl acceptor.

Our experiments show that the two enzymes exhibit many similarities in nucleotide binding and in metal ion requirements. There are, however, some distinctions. Phosphotransferase activity with the cAMP-dependent enzyme with lin-benzo-ATP as donor is very poor in contrast to that of the cAMP-dependent enzyme. The preferred synthetic acceptor peptide substrate of the cGMP-dependent enzyme decreases nucleotide affinity; no such effect is observed with the cAMP-dependent enzyme.

**Materials and Methods**

The synthetic heptapeptide (Ser-peptide) used as substrate (Leu-Arg-Arg-Ala-Ser-Leu-Gly) was purchased from Boehringer Mannheim Biochemicals, and Arg-Lys-Arg-Ser-Arg-Ala-Glu ([Ala$^{34}$]histone H2B-(29-35)) was synthesized as described by Glass & Krebs (1982). Carrier-free [γ-32P]ATP and inorganic [32P]phosphate were purchased from ICN, lin-benzo-[γ-32P]ATP was synthesized from 20-25 nmol of lin-benzo-ADP and 15 mCi of inorganic [32P]phosphate by the method of Walseth & Johnson (1979), and the product was purified as described for [γ-32P]ATP by Palmer & Avruch (1981). The salts of metal ions were purchased from Aldrich Chemical Co. All other chemicals were purchased from Sigma Chemical Co. The concentrations of lin-benzoadenine nucleotides were measured by absorbance at 331 nm by using an extinction coefficient of 9750 M$^{-1}$ cm$^{-1}$ (Leonard et al., 1976).

Protein Kinase Preparations and Activity Measurements. The cGMP-dependent protein kinase was purified from bovine lung as described by Glass & Krebs (1979). The preparation had a specific activity of 5.5 $\mu$mol (min·mg$^{-1}$) by using 0.5 mg/mL histone H2B as substrate under the conditions described by Glass & Krebs (1979). The activity ratio of these preparations of the enzyme using 25 $\mu$M histone H2B-(29-35) as substrate was 0.09, indicating an 11-fold stimulation of activity by cGMP.

The type II catalytic subunit of the cAMP-dependent protein kinase from bovine heart was purified as described by Hartl & Roskoski (1982). Phosphotransferase activity measurements of the catalytic subunit were performed as described by Hartl & Roskoski (1982) and Roskoski (1983). Protein concentrations were determined by the procedures of either Lowry et al. (1951) or Bradford (1976) using bovine serum albumin or ovalbumin as standards, respectively. For the determination of the molarity of the enzymes, the molecular weight of the catalytic subunit of the cAMP-dependent protein kinase was taken to be 40 000 and that of the cGMP-dependent protein kinase as 154 000.

**Steady-State Kinetic Assays.** Phosphotransferase activity of cGMP-dependent protein kinase was measured at 30 °C by using the phosphocellulose paper adsorption method of Roskoski (1983). lin-Benzo-[γ-32P]ATP and [γ-32P]ATP (10-100 $\mu$M) were used as the phosphoryl donors with specific activities of 100-200 and 40-150 cpm/pmol, respectively. Ser-peptide (1400 $\mu$M) or [Ala$^{34}$]histone H2B-(29-35) (200 $\mu$M) were used as the phosphoryl acceptors. The final volume (80 $\mu$L) of the reaction mixture contained 50 mM Mops, pH 7.0, 2 $\mu$M cGMP, 10 mM MgSO$_4$, 0.3 mg/mL bovine serum albumin, and 3 mM 2-mercaptoethanol. Assays were conducted with the amounts of cGMP-dependent protein kinase and for the times indicated in the tables and figure legends. These conditions were chosen on the basis of the substrates being used. Reactions were linear with respect to time and amount of enzyme. All values are the means and standard errors of three or more determinations.

**Fluorescence Polarization Titrations.** The fluorescence measurements were performed with a SLM4800 spectrofluorometer interfaced with a Hewlett-Packard HP 9825 A calculator. Fluorescence polarization was calculated by using a program supplied by SLM Instruments Inc. Polarization (P) and anisotropy ($r$) are defined as

$$P = \frac{I_p - I_\perp}{I_p + 2I_\perp}$$

$$r = \frac{I_p - I_\perp}{I_p + 2I_\perp}$$

where, $I_p$ and $I_\perp$ are the intensities observed parallel and perpendicular to the polarization of the exciting light, respectively. Polarization and anisotropy values of the lin-benzoadenine nucleotides were determined by using calcite polarizers. Excitation was at 334 nm, with 4-nm resolution, and emitted light was isolated with a Schott KV 389 filter.

Three types of polarization titrations were performed in order to determine the binding constants of various nucleotides for the cGMP-dependent protein kinase. The titrations were performed in 50 mM Mops (pH 7.0) and 100 mM NaCl in the presence or absence of 20 $\mu$M cGMP. This concentration of cGMP was sufficient to saturate the protein kinase even under the conditions of high concentration of the enzyme used in the fluorescence polarization studies. All experimental values represent the means and standard errors of at least three independent determinations.

**Dilution Titration.** Polarization as a function of varying enzyme concentration (at constant [lin-benzoadenine nucleotide]) was measured as described in our earlier reports (Hartl et al., 1983; Bhatnagar et al., 1983) to determine $P_0$. $P_0$ is the polarization value when all lin-benzoadenine nucleotide is bound to the protein kinase (at infinite enzyme concentration).

The theoretical polarization ($P_0$) of lin-benzo-ADP rigidly bound to the protein kinase and the average angle of rotation ($\omega$) of rigidly bound lin-benzo-ADP to the enzyme were calculated from Perrin’s equation (Perrin, 1926) as described earlier for the cAMP-dependent enzyme (Hartl et al., 1983). The $P_0$, the anisotropy value when all lin-benzoadenine nucleotide is bound to the enzyme, was determined from anisotropy data obtained simultaneously. The theoretical anisotropy ($r_0$) of lin-benzo-ADP rigidly bound to the protein kinase was calculated from the following Perrin’s equation (Lakowicz, 1983):

$$\frac{1}{r_0} = \frac{1}{r_0} \left(1 + \frac{RT\tau}{\eta V_0}\right)$$

where $r_0$ is the intrinsic anisotropy of the fluorophore, $R$ is the universal gas constant (erg·mol$^{-1}$·deg$^{-1}$), $T$ is the absolute temperature (K), $\eta$ is the viscosity (centipoise), $\tau$ is the lifetime of the excited state ($\tau \approx 4.2$ ns for lin-benzo-ADP; VanDerLijn et al., 1978a), and $V_0$ is the molecular volume of the fluorescent unit (cm$^3$/g).

The value of $r_0$ for lin-benzo-ADP in the above equation was determined by measuring the fluorescence anisotropy in solutions of varying viscosity with sucrose or glycerol; $1/\tau$ was plotted vs. $T/\eta$, and $1/r_0$ was obtained from the ordinate intercept where $\eta \to \infty$. The value of $r_0$ was calculated to be
0.301 for linbenzo-ADP at 334 nm. The average angle of rotation (ωo) of linbenzo-ADP rigidly bound to the cGMP-dependent protein kinase was calculated from anisotropy (r0) data according to the following equation (Lakowicz, 1983):

$$\frac{1}{r_0} - 1 = \frac{3}{r_0} (\cos^2 \omega_0 - 1)$$

(b) Addition Titration. Polarization (P_{obs}) was recorded as described previously (Hartl et al., 1983; Bhatnagar et al., 1983) at each nucleotide concentration after addition of successive increments of linbenzoadenine nucleotide to a constant protein kinase concentration. The dissociation constant (Kd) of linbenzoadenine nucleotide for the enzyme was then determined by Scatchard analysis (Scatchard, 1949).

(c) Displacement Titration. The dissociation constant (Kd) of ADP for the cGMP-dependent protein kinase was determined by displacing the fluorescent linbenzo-ADP bound to the enzyme with ADP as previously described for other nucleotides (Bhatnagar et al., 1983).

RESULTS

Interaction of H2B-(29-35) with cGMP-Dependent Protein Kinase. The enzyme velocity was determined with varying ATP concentrations (10-100 µM) at a fixed, near saturating [Ala34]histone H2B-(29-35) concentration (200 µM; Glass & Krebs, 1982) in the presence of various fixed concentrations of linbenzo-ADP (0.8, 16.6, 24.9, and 33.2 µM). Mg-linbenzo-ADP was a competitive inhibitor with respect to MgATP (not shown). A slope vs. [linbenzo-ADP] replot was linear and gave a Kd for linbenzo-ADP of 21.5 ± 0.35 µM. The Kd for ADP obtained in parallel experiments was 169 ± 54 pM (Table I). Thus, linbenzo-ADP, which is 2.4 Å wider in the adenine moiety, interacts with the active site of the protein kinase with a higher affinity than does the natural reaction product, ADP.

[Ala34]histone H2B-(29-35) was selected as the phosphate acceptor for the cGMP-dependent protein kinase in these studies since it contains only one site of phosphorylation (as compared to various histones) and is one of the best known substrates (low Km and high V_{max} for the enzyme (Glass et al., 1981; Glass & Krebs, 1982). The peptide is an analogue of the sequence around serine-32 in histone H2B in which an alanine residue has been substituted for the lysine normally present at position 34. Both peptides have similar Km values (20-30 µM), but the Ala34 replacement analogue has a 4-5-fold higher V_{max} (20 µmol (min-mg)^{-1}) than that of the parent peptide, histone H2B-(29-35), at pH 7.4 in Tris buffer (Glass & Krebs, 1982). The pH optimum (Figure 1) for the phosphorylation of histone H2B-(29-35) is rather high [see also Glass & Krebs (1979)]. One possible reason for this is that the lysine in position 34 is a negative determinant (Glass & Krebs, 1982) when in the protonated form (Glass, unpublished results). The pH profile of [Ala34]histone H2B-(29-35), which lacks lysine-34, does not, therefore, show this behavior (Figure 1). Since we wished to perform our studies in Mops buffer at pH 7.0 (for comparison with our earlier studies with the cAMP-dependent enzyme), [Ala34]histone H2B-(29-35) was the more suitable substrate. Ser-peptide was not routinely used for the steady-state kinetic studies because, even though the cGMP-dependent protein kinase has a high V_{max} with Serpeptide, its K_{m} is high as well (approximately 200 µM; Glass & Krebs, 1979; Glass et al., 1981). This meant that we would have had to use Ser-peptide at a concentration of well over 1 mM to even approach saturation.

Measurement of the K_{d} for linbenzo-ADP and ADP by Fluorescence Polarization and Anisotropy Titrations. For characterization of the interaction of linbenzo-ADP with the cGMP-dependent protein kinase in the presence and absence of cGMP, fluorescence polarization titrations were performed. linbenzo-ADP was used instead of linbenzo-ATP because the protein kinase demonstrated intrinsic ATPase activity (Glass and O'Neill, unpublished observations) for which linbenzo-ATP was a substrate (not shown). No difference was observed in the fluorescence emission spectrum of linbenzo-ADP in the presence or absence of the protein kinase with or without saturating cGMP (not shown). Binding of the nucleotide to the enzyme, however, was accompanied by an increase in polarization. The binding constants of linbenzo-ADP for the cGMP-dependent protein kinase were measured by fluorescence polarization dilution and addition titrations and those of ADP by fluorescence polarization displacement titrations as described under Materials and Methods.

The polarization (P_{obs}) of linbenzo-ADP bound to the protein kinase was determined by dilution titration (Figure 2A). At constant linbenzo-ADP concentration in the presence of 10 mM Mg^{2+}, the fluorescence-polarization (P_{obs}) increases with increasing enzyme concentration as more ligand is bound. P_{obs} ranges from P_{0} (zero protein concentration) to P_{0} (all ligand

![Figure 1: Effect of pH on the phosphorylation of histone H2B-(29-35) (A) and [Ala34]histone H2B-(29-35) (B) by cGMP-dependent protein kinase. Both synthetic peptides were used at a concentration of 250 µM. The enzyme was used at a concentration of 0.5 µg/mL. Reaction velocities were determined under standard assay conditions as described under Materials and Methods except that the pH was changed. Reactions were maintained at the indicated pH with the appropriately adjusted MES/Tris (50 mM/50 mM) buffers. Reactions were conducted for 5 min.](https://example.com/figure1.png)
is bound to the protein). Extrapolation of a plot of (polarization) vs. [protein kinase] to zero resulted in a value of $P_b$ of nearly 0.35 (Figure 3). This corresponds well to a $P_b$ value of 0.370 calculated by the Perrin equation (Perrin, 1926) for rigidly bound $lin$-benzo-ADP (Table II). The values of $P_b$ obtained in the absence or presence of 20 mM GMP were similar (Table II). The observed $P_b$ was used to calculate the average angle of rotation ($\omega_p$) of $lin$-benzo-ADP bound to the cGMP-dependent protein kinase, in the absence or presence of 20 mM GMP, during the lifetime of its excited state (4.2 ns; VanDerLijn et al., 1978a). The values of 17° and 18° for $\omega_p$ obtained in either the absence or presence of cGMP, respectively, are very close to the theoretical value of 14° calculated for the rotation of the protein from the Perrin equation. $P_b$, the intrinsic polarization of the fluorophore, and $P_{bound}$, the polarization in the bound state, are similar within experimental error. Thus, most or all of the observed rotation of $lin$-benzo-ADP bound to the enzyme is due to the rotation of the protein itself. This also indicates that $lin$-benzo-ADP binds rigidly to the cGMP-dependent protein kinase in both the absence or presence of cGMP, so that there is little or no rotation within the adenine binding site.

Anisotropy values were also calculated (by the definition given under Materials and Methods) from the fluorescence intensity data obtained during the titration. The values of $r_0$.

![Figure 2: Polarization titrations of the cGMP-dependent protein kinase with $lin$-benzo-ADP. Measurements were performed at 23 °C in 50 mM Mops (pH 7.0), 100 mM NaCl, and 10 mM MgSO4. (Panel A) Dilution titration. The protein kinase was varied (17.5-3.8 pM) at a constant (2 nM) $lin$-benzo-ADP concentration in the presence of 10 mM MgSO4 and 20 pM cGMP. (Panel B) Addition titration. $lin$-Benzo-ADP was varied at a constant (3 nM) protein kinase concentration in 150-pL total volume by addition of 1-pL portions of a concentrated $lin$-benzo-ADP solution. The titration shown was performed in the presence of 20 pM cGMP and 200 pM phosphor-rylatable peptide [Ala24]histone H2B-(29-35). Corrections for the changes in volume and ligand concentration were routinely made.](image2)

![Figure 3: Determination of fluorescence polarization ($P_b$) and anisotropy ($r_0$) for the protein kinase. Dilution titrations were performed with the protein kinase in the presence of 20 pM cGMP as described in the legend to Figure 2 and under Materials and Methods. From the ordinate intercepts of a plot of (polarization)$^2$ (A) or (anisotropy)$^{-1}$ (B) vs. [protein kinase]$^{-1}$, a $P_b$ value of 0.350 and an $r_0$ value of 0.264 were obtained for the protein kinase in the presence of 20 pM cGMP.](image3)
Table III: Kinetic Parameters of cGMP-Dependent Protein Kinase
with ATP and lin-benzo-ATP as Substrates

<table>
<thead>
<tr>
<th>substrate</th>
<th>$K_m$ (µM)</th>
<th>$V_{max}$ [µmol (min-ng)$^{-1}$]</th>
</tr>
</thead>
<tbody>
<tr>
<td>ATP</td>
<td>32.2 ± 0.57</td>
<td>11.1 ± 0.17</td>
</tr>
<tr>
<td>lin-benzo-ATP</td>
<td>28.5 ± 2.3</td>
<td>0.00623 ± 0.00035</td>
</tr>
</tbody>
</table>

* Determined by radioisotopic assay with [Ala$^{34}$] histone H2B-(29-35) as phosphoryl acceptor as described under Materials and Methods.

Reactions were performed at a fixed, near saturating concentration of the peptide substrate (200 µM) and Mg$^{2+}$ concentration of 10 mM. Nucleotide concentrations were varied from 10 to 100 µM. When ATP was the substrate, assays were conducted for 2 min at a protein kinase concentration of 0.3–0.4 µg/mL. When lin-benzo-ATP was the substrate, assays were conducted for 10–15 min with 8–15 µg/mL enzyme. $K_m$ and $V_{max}$ were determined both from double-reciprocal plots ($v_0$ vs. [nucleotide]$^-1$) and from plots of $v_0$ vs. $v_0$/[nucleotide] according to Hofstee (1952).

I) with a stoichiometry of 2 mol of lin-benzo-ADP bound per mol of the enzyme dimer. The $K_d$ value of lin-benzo-ADP, however, is 2-fold greater than the $K_d$ value of ATP (Table I). The addition of this peptide substrate to the fluorescence titration decreases the binding affinity of lin-benzo-ADP for the enzyme 2-fold (from 11 to 23 µM) while the number of binding sites remain constant (2 mol/mole of enzyme dimer). The $K_d$ of lin-benzo-ADP of 23 µM determined in the presence of [Ala$^{34}$]histone H2B-(29-35) then corresponds well with the $K_d$ for lin-benzo-ADP of 22 µM. The $K_d$ of lin-benzo-ADP is in good agreement with $K_d$ of ADP for the enzyme when both $K_d$'s are obtained in the absence of the peptide substrate (Table I). But in the presence of [Ala$^{34}$]histone H2B-(29-35) the $K_d$ of ADP for the enzyme is 5-fold higher than that of lin-benzo-ADP and is in agreement with the $K_d$ of 169 µM of ADP for the cGMP-dependent protein kinase (Table I).

The $K_d$ values of lin-benzo-ADP and ADP under various conditions were calculated on the basis of anisotropy as well. The maximum difference between these values obtained from anisotropy and polarization data was within experimental error (less than 2%). The determinations in this study have been based on polarization values to maintain continuity with earlier work (Hartl et al., 1983; Bhatnagar et al., 1983).

lin-Benzo-ATP as a Substrate for cGMP-Dependent Protein Kinase. To determine the effectiveness of lin-benzo-ATP as a phosphoryl donor for the reaction catalyzed by the cGMP-dependent protein kinase, steady-state kinetic studies were performed with lin-benzo-ATP or ATP and [Ala$^{34}$]histone H2B-(29-35) as phosphoryl acceptor. lin-Benzo-ATP has a $K_m$ of 29 µM which corresponds well with the $K_m$ for ATP of 32 µM (Table III). However, the $V_{max}$ with lin-benzo-ATP is over 3 orders of magnitude less than that with ATP as substrate (Table III). Similar results were observed when Ser-peptide was used as the phosphoryl acceptor.

Metal Ion Dependence of Nucleotide Binding and Phosphotransferase Activity. As mentioned earlier, nucleotide binding to the protein kinase is dependent on the presence of a divalent cation. On addition of metal ion, lin-benzo-ADP binds to the enzyme, resulting in an increase in polarization. With increasing Ca$^{2+}$ concentration, for example, the fluorescence polarization is enhanced, which indicates that an increasing amount of lin-benzo-ADP is bound to the enzyme (Figure 4). The binding of the nucleotide to the cGMP-dependent protein kinase is half-maximum at 1.0 mM Ca$^{2+}$ and plateaus at approximately 10 mM. The $K_d$ of 1.2 mM for Ca$^{2+}$ was determined from a plot of (polarization)$^-1$ vs. [Ca$^{2+}$]$^-1$ (Figure 4, inset).

![Figure 4: Metal ion requirement of nucleotide binding to the cGMP-dependent protein kinase.](image)

Table IV: Dependence of Nucleotide Binding and Phosphotransferase Activity of cGMP-Dependent Protein Kinase on Metal Ions

<table>
<thead>
<tr>
<th>metal ion</th>
<th>$K_d$ (µM)</th>
<th>phosphotransferase activity (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>none</td>
<td>ND</td>
<td>&lt;0.6</td>
</tr>
<tr>
<td>Mg$^{2+}$</td>
<td>1.4 ± 0.18</td>
<td>100.0</td>
</tr>
<tr>
<td>Mn$^{2+}$</td>
<td>0.144 ± 0.008</td>
<td>6.5 ± 0.1</td>
</tr>
<tr>
<td>Co$^{2+}$</td>
<td>0.723 ± 0.012</td>
<td>23.7 ± 0.6</td>
</tr>
<tr>
<td>Cd$^{2+}$</td>
<td>ND</td>
<td>&lt;0.2</td>
</tr>
<tr>
<td>Ca$^{2+}$</td>
<td>1.2 ± 0.10</td>
<td>&lt;0.5</td>
</tr>
<tr>
<td>Sr$^{2+}$</td>
<td>4.2 ± 0.82</td>
<td>&lt;0.3</td>
</tr>
<tr>
<td>Ba$^{2+}$</td>
<td>4.6 ± 0.66</td>
<td>&lt;0.6</td>
</tr>
<tr>
<td>Fe$^{2+}$</td>
<td>ND</td>
<td>ND</td>
</tr>
<tr>
<td>Ni$^{2+}$</td>
<td>1.6 ± 0.12</td>
<td>30.4 ± 0.9</td>
</tr>
<tr>
<td>Zn$^{2+}$</td>
<td>ND</td>
<td>&lt;0.2</td>
</tr>
<tr>
<td>Cu$^{2+}$</td>
<td>ND</td>
<td>ND</td>
</tr>
<tr>
<td>Gd$^{3+}$</td>
<td>ND</td>
<td>ND</td>
</tr>
<tr>
<td>La$^{3+}$</td>
<td>ND</td>
<td>ND</td>
</tr>
</tbody>
</table>

* Determined by fluorescence polarization enhancement procedure as described in the legend of Figure 4. For catalytic activity determinations, all metal ions were added as their CI$^-$ salts (except for Cd$^{2+}$, which was added as CdSO$_4$). The metal ions that supported nucleotide binding were used at final concentrations of 3 times their apparent $K_d$ values. Metal ions that did not support binding were used at a concentration of 10 mM. Activities are expressed as a percentage of the activity in the presence of 4.2 mM MgCl$_2$, which corresponds to a catalytic activity of 14.2 ± 0.7 µmol (min-ng)$^{-1}$ (n = 7). The activity measurements were made in the presence of 400 µM [γ-32P]ATP and 1400 µM Ser-peptide with 1.2 µg/mL protein kinase for 2 min. ND, not detectable.

A number of metal ions (Mg$^{2+}$, Mn$^{2+}$, Co$^{2+}$, Ca$^{2+}$, Sr$^{2+}$, Ba$^{2+}$, and Ni$^{2+}$) promoted nucleotide binding to the enzyme as demonstrated by an enhancement in fluorescence polarization (Table IV). There was no increase in fluorescence polarization caused by these metal ions in the presence of lin-benzo-ADP without the enzyme. On the other hand, Cd$^{2+}$, Fe$^{2+}$, Zn$^{2+}$, Cu$^{2+}$, Gd$^{3+}$, and La$^{3+}$ failed to promote any binding under our experimental conditions. Cd$^{2+}$, moreover, seemed to denature and precipitate the enzyme since addition of 5 mM Mg$^{2+}$, following a Cd$^{2+}$ titration, failed to promote nucleotide binding.

Subsaturating metal ion concentrations (3 times the $K_d$ of metal ions) were used to determine which metals support...
phosphotransferase activity. Under these conditions, the catalytic activity was maximal \([14.2 \, \mu\text{mL}^{-1} \cdot \text{min}^{-1}]\) in the presence of 4.2 mM \(\text{Mg}^{2+}\) (Table IV). Of the other metal ions tested \(\text{Ni}^{2+}\) and \(\text{Co}^{2+}\) were most effective. At a 2.2 mM concentration, \(\text{Co}^{2+}\) sustained a catalytic activity of 3.37 \(\mu\text{mol} \cdot \text{min}^{-1}\); with 4.8 mM \(\text{Ni}^{2+}\), an activity of 4.32 \(\mu\text{mol} \cdot \text{min}^{-1}\) was recorded. \(\text{Mn}^{2+}\) (432 \(\mu\text{M}\)) supported less than 7% of the activity of the enzyme as compared to that supported by 4.2 mM \(\text{Mg}^{2+}\). Although \(\text{Ca}^{2+}\), \(\text{Sr}^{2+}\), and \(\text{Ba}^{2+}\) promoted binding of the nucleotide to the enzyme (Table IV), no significant phosphotransferase activity of the cGMP-dependent enzyme was measured in the presence of any of these metal ions at concentrations 3 times their respective \(K_m\) values under our experimental conditions. \(\text{Cd}^{2+}\), \(\text{Fe}^{2+}\), \(\text{Zn}^{2+}\), \(\text{Cu}^{2+}\), \(\text{Gd}^{3+}\), and \(\text{La}^{3+}\) at concentrations of 10 mM failed to support detectable phosphotransferase activity.

The cGMP-dependent protein kinase is similar to the cAMP-dependent enzyme in that increasing the concentration of \(\text{Mg}^{2+}\) lowers the \(K_m\) for ATP but also lowers the \(V_{\max}\) of the reaction (Glass & Krebs, 1979; Cook et al., 1982). For example, in our experiments with 100 \(\mu\text{M}\) \([\text{Ala}^{34}]\text{histone H2B-(29-35)}\) (Table I), the \(K_m\) for ATP was 44.9 \(\mu\text{M}\) and the \(V_{\max}\) was 13.6 \(\mu\text{mol} \cdot \text{min}^{-1}\) as compared with the \(K_m\) of ATP of 32.2 \(\mu\text{M}\) and \(V_{\max}\) of 11.1 \(\mu\text{mol} \cdot \text{min}^{-1}\) at 10 mM \(\text{Mg}^{2+}\). The effect on the cAMP-dependent enzyme, however, was more dramatic.

**DISCUSSION**

We have studied the interaction of \(\text{lin}-\text{benzoadenine nucleotides with the cGMP-dependent protein kinase.}\ lin-Benzo-ADP, which is stretched out by 2.4 \(\AA\) over ADP in its adenine moiety (Scopes et al., 1977), is a linear competitive inhibitor (Cleland, 1970) with respect to Mg-ATP as is ADP (as described under Results). In addition, the \(K_m\) for \(\text{lin}-\text{benzo-ADP}\) determined by fluorescence polarization studies (in the absence of peptide substrate) is very close to the \(K_m\) determined for ADP under similar conditions (Table I). Thus, \(\text{lin}-\text{benzo-ADP}\) binds to the active site of the enzyme as does ADP. However, \(\text{lin}-\text{benzo-ADP}\) binds to the protein kinase [in the presence of peptide substrate \([\text{Ala}^{34}]\text{histone H2B-(29-35)}\)] with over 6-fold tighter affinity than does ADP (Table I). For the cAMP-dependent protein kinase catalytic subunit, on the other hand, the \(K_m\) of \(\text{lin}-\text{benzo-ADP}\) determined by fluorescence polarization studies was found to be very close to the kinetically determined \(K_m\) of ADP (Hartl et al., 1983) with Ser-peptide as substrate. Unlike the cGMP-dependent enzyme, the \(K_m\) of \(\text{lin}-\text{benzo-ADP}\) for the cAMP-dependent protein kinase catalytic subunit was not altered by 200 \(\mu\text{M}\) \([\text{Ala}^{34}]\text{histone H2B-(29-35)}\) with over 6-fold tighter affinity than does ADP (Table I). For the cAMP-dependent protein kinase catalytic subunit, on the other hand, the \(K_m\) of \(\text{lin}-\text{benzo-ADP}\) (from inhibition studies) and its \(K_m\) (from fluorescence polarization studies) were found to be very close to the kinetically determined \(K_m\) for ADP (Hartl et al., 1983) with Ser-peptide as substrate. Unlike the cGMP-dependent enzyme, the \(K_m\) of \(\text{lin}-\text{benzo-ADP}\) for the cAMP-dependent protein kinase catalytic subunit was not altered by 200 \(\mu\text{M}\) \([\text{Ala}^{34}]\text{histone H2B-(29-35)}\) \((K_m = 10.2 \, \mu\text{M}\) in the absence of peptide and 9.8 \(\mu\text{M}\) in the presence of peptide). These \(K_m\) values of \(\text{lin}-\text{benzo-ADP}\) for the cAMP-dependent enzyme correspond well with the \(K_m\) of 8.9 \(\mu\text{M}\) of ADP in the presence of 200 \(\mu\text{M}\) \([\text{Ala}^{34}]\text{histone H2B-(29-35)}\).

The polarization data in Table II indicate that the \(\text{lin}-\text{benzoadenine moiety is rigidly bound to the cGMP-dependent protein kinase within the limits of detection. This parallels our observations for the cAMP-dependent protein kinase catalytic subunit (Hartl et al., 1983; Bhatnagar et al., 1983). Granot et al. (1979) showed by NMR that there is a strong interaction between the catalytic subunit of cAMP-dependent protein kinase and the adenosine portion of Co(NH$_3$)$_2$ATP. This differs from the binding of \(\text{lin}-\text{benzo-ATP}\) to \(E.\text{coli}\) aspartate transcarbamylase (Chien & Weber, 1973; VanDerLijn et al., 1978b) and pyruvate kinase (Barrio et al., 1973). In the latter two cases, \(\text{lin}-\text{benzo-ATP}\) is bound in a manner such that there is appreciable rotational freedom within the nucleotide binding site as demonstrated by fluorescence polarization.

As summarized in Table I, there is a significant effect of the presence of the peptide substrate, \([\text{Ala}^{34}]\text{histone H2B-(29-35)}\), on the binding affinity of both ADP and \(\text{lin}-\text{benzo-ADP}\) to the cGMP-dependent protein kinase, but this is not so for the cAMP-dependent protein kinase (not shown). In addition, \(\text{lin}-\text{benzo-ADP}\) binds to the cGMP-dependent enzyme with greater affinity than does ADP (Table I); \(\text{lin}-\text{benzo-ADP}\) and ADP, on the other hand, bind to the catalytic subunit of the cAMP-dependent protein kinase with essentially equal affinities (Hartl et al., 1983; Bhatnagar et al., 1983). The large difference between the \(K_m\) values for \(\text{lin}-\text{benzo-ADP}\) and ADP itself (for the cGMP-dependent enzyme; Table I) was due not merely to the use of \([\text{Ala}^{34}]\text{histone H2B-(29-35)}\) as the phosphate-accepting substrate. When Ser-peptide (1400 \(\mu\text{M}\)) was used as the substrate in the presence of 10 mM \(\text{Mg}^{2+}\), the \(K_m\) for \(\text{lin}-\text{benzo-ADP}\) was 15.7 \(\mu\text{M}\) and the \(K_m\) for ADP was 132 \(\mu\text{M}\). The \(K_m\) of ADP of 169 \(\mu\text{M}\), moreover, is similar to the value of 265 \(\mu\text{M}\) previously reported (Glass et al., 1981) when a different buffer and pH were used. Even at lower \(\text{Mg}^{2+}\) (3 mM), the \(K_m\) of \(\text{lin}-\text{benzo-ADP}\) was 17.7 \(\mu\text{M}\) and that of ADP was 233 \(\mu\text{M}\) when \([\text{Ala}^{34}]\text{histone H2B-(29-35)}\) was used as the substrate.

The decrease in binding affinity of adenine nucleotides due to the peptide could be because \([\text{Ala}^{34}]\text{histone H2B-(29-35)}\) is so positively charged that it interacts ionically with either the adenine nucleotide or the enzyme in such a way so as to disturb the binding of the nucleotide to the active site of the enzyme. In any event, the conclusion that the peptide decreases the affinity of the enzyme for adenine nucleotide is consistent with earlier reports of the pattern of the initial velocity kinetics (Glass et al., 1981). Steady-state kinetic studies using the parent peptide histone H2B-(29-35) as the acceptor and ATP as the donor show that the lines intersect at a point below the x axis (Figure 5). As the concentration of the histone peptide is increased, the apparent \(K_m\) value of ATP is increased (i.e., a decreased affinity). Varying the concentration of ATP affects the apparent \(K_m\) values of histone peptide in a similar fashion (data not shown). This type of initial velocity plot is not seen with all peptide substrates of the cGMP-dependent protein kinase. Specifically, the kinetics for ATP and histone H2B-(29-35) as substrates. The enzyme (1.0 \(\mu\text{g/mL}\)) was incubated for 2 min at 30 \(^\circ\text{C}\) with \([\gamma-32\text{P}]\text{ATP}\) concentrations from 10 to 100 \(\mu\text{M}\) at fixed histone H2B-(29-35) concentrations of 10 (●), 15 (△), 25 (○), and 50 (●) \(\mu\text{M}\). Activity was measured as described under Materials and Methods except the buffer was 30 mM Tris-HCl (pH 7.4) and the \(\text{MgSO}_4\) concentration was 3 mM. (Inset) Replot of the velocity$^{-1}$-axis intercept vs. [peptide]$^{-1}$.
performed with ATP and a peptide corresponding to the autophosphorylation site in the enzyme (Glass & Smith, 1983) yield initial velocity lines that intersect above the x axis (Glass, unpublished observations). In this case, as the concentration of one substrate is increased, the apparent $K_m$ of the other substrate is decreased.

The large difference in the affinities of cGMP-dependent protein kinase for lin-benzo-ADP as compared to ADP (Table I) in the presence of [Ala$^{2+}$]histone H2B-(29-35) might be due to a conformational change induced by the peptide. A comparable phenomenon has been extensively studied with hexokinase (Bennett & Steitz, 1980) where glucose binds to the enzyme prior to ATP and brings about a conformational change of the two lobes of the hexokinase molecule so as to place the ATP molecule in a suitable position for the phosphorylation of glucose. In the case of the cGMP-dependent protein kinase, the $K_m$ values are essentially the same with ATP or lin-benzo-ATP as substrate, but the $V_{\text{max}}$ with lin-benzo-ATP is very low, over 3 orders of magnitude less than that with ATP. This also suggests that in the presence of the peptide the $\gamma$-phosphate of lin-benzo-ATP is not in a suitable position for phosphotransferase activity to occur possibly due to the lateral stretching of 2.4 A of the adenine ring system (in lin-benzo nucleotides). Most enzymes that utilize lin-benzo-ATP as substrate exhibit a lower $V_{\text{max}}$ with a $K_m$ comparable to that seen with ATP (Leonard et al., 1978). Alternative mechanisms, however, might be responsible for the low $V_{\text{max}}$ seen with the cGMP-dependent protein kinase. To assure that the low $V_{\text{max}}$ values were not due to technical problems with the lin-benzo[32P]ATP preparations, they were tested with the catalytic subunit of the cAMP-dependent enzyme. The $K_m$ of catalytic subunit for lin-benzo-ATP was 14.9 $\mu$M, and the $V_{\text{max}}$ was 4.03 $\mu$mol (min-mg)$^{-1}$ with the lin-benzo-[32P]ATP synthesized for this study. These values are in good agreement with our previous results [11.3 $\mu$M for $K_m$ and 5.0 $\mu$mol (min-mg)$^{-1}$; Hartl et al., 1983].

The $K_i$'s of lin-benzo-ADP or ADP for the catalytic subunit of the cAMP-dependent protein kinase are not altered either in the presence of [Ala$^{2+}$]histone H2B-(29-35) (as stated earlier) or with Ser-peptide (Hartl et al., 1983; Bhatnagar et al., 1983). With that enzyme lin-benzo-ATP, lin-benzo-ADP, ATP, and ADP have the same affinity for the enzyme (Hartl et al., 1983; Bhatnagar et al., 1983). Also, lin-benzo-ATP is a good substrate for the catalytic subunit, with a $K_m$ equal to that of ATP and a $V_{\text{max}}$ 20% that of the $V_{\text{max}}$ of ATP. Therefore, unlike the cGMP-dependent enzyme, the cAMP-dependent protein kinase appears to accommodate lin-benzo-ADP and lin-benzo-ATP more like ADP and ATP, respectively, both in the absence or in the presence of the peptide substrate.

The requirement of a divalent metal cation for the enzyme activity of the cGMP-dependent protein kinase is well established (Kuo et al., 1978). As stated under Results, binding of lin-benzo-ADP is enhanced (increased fluorescence polarization) by increasing the concentration of metal ion. This further shows that the metal ion is required for nucleotide binding to the cGMP-dependent enzyme. We suggest that the apparent $K_m$ values (Table IV) reflect the affinities of the metal ions for an inhibitory binding site on the enzyme similar to that discussed earlier for cAMP-dependent protein kinase catalytic subunit (Granot et al., 1979; Bhatnagar et al., 1983). The rank order of metal ion effectiveness in supporting phosphotransferase activity was $\text{Mg}^{2+} > \text{Ni}^{2+} > \text{Co}^{2+} > \text{Mn}^{2+}$. To our knowledge this is the first report as to the ability of $\text{Ni}^{2+}$ to support nucleotide binding and catalysis of the cGMP-dependent protein kinase. Otherwise, these results are generally consistent with most of the early reports on the mammalian cGMP-dependent protein kinase: i.e., $\text{Co}^{2+}$ is the next best metal after $\text{Mg}^{2+}$ in supporting catalysis, and $\text{Mn}^{2+}$ will do so only weakly (Hofmann & Sold, 1972; Nakazawa & Sano, 1975; Kuo et al., 1976; Shoji et al., 1977a,b; Gill et al., 1977). In an early report with the arthropod enzyme, however, the rank order was $\text{Co}^{2+} > \text{Mn}^{2+} > \text{Mg}^{2+}$ in supporting catalysis (Kuo & Greengard, 1970). More recently, Hixson & Krebs (1981) have reported that $\text{Co}^{2+}$ increases the pseudo-first-order rate constant of inactivation of the cGMP-dependent protein kinase by $\text{S}^2$-[-(fluorosulfonyl)]benzoyladenosine compared to the rate constant in the absence of metal ion; $\text{Co}^{2+}$ was better than $\text{Mg}^{2+}$ in doing so. No correlation between the ionic radii or the electronic configurations of the metal ions could be established with either nucleotide binding or catalytic activity.

The metal ion requirement of the cGMP-dependent protein kinase, apart from the actual $K_i$ values of metal ions for the enzymes, differs from that of the catalytic subunit of the cAMP-dependent enzyme in the effectiveness of the metal ions in supporting catalysis. The rank order for the catalytic subunit of the cAMP-dependent enzyme was $\text{Mg}^{2+} > \text{Mn}^{2+} > \text{Co}^{2+} > \text{Cd}^{2+}$ (Bhatnagar et al., 1983). $\text{Ni}^{2+}$ and $\text{Ba}^{2+}$ did not support either catalysis or nucleotide binding in the case of the cAMP-dependent protein kinase (Bhatnagar et al., 1983), whereas $\text{Ni}^{2+}$ supported both catalysis and nucleotide binding and $\text{Ba}^{2+}$ only nucleotide binding to the cGMP-dependent protein kinase (Table IV). $\text{Cd}^{2+}$, on the other hand, did not support either catalysis or nucleotide binding with the cGMP-dependent enzyme but was effective for both functions with the catalytic subunit of the cAMP-dependent enzyme.

In conclusion, the interactions of lin-benzoadenine di- and triphosphates with the cGMP-dependent protein kinase (in this study) were similar for most part to those observed with the cAMP-dependent protein kinase (Hartl et al., 1983; Bhatnagar et al., 1983). The major differences were in the $V_{\text{max}}$ for lin-benzo-ATP and the effect of peptide substrates on nucleotide binding. There appears to be some conformational change in the cGMP-dependent protein kinase on peptide substrate binding which seems to distort the nucleotide binding site, especially the adenine binding region. The determinants within the histone peptide producing such an effect might be conveniently studied by determining the influences of a series of synthetic peptide analogues (Glass & Krebs, 1982) on the binding of lin-benzo-ADP. This conformational change may be responsible for the extremely low $V_{\text{max}}$ (almost one-two thousandth of that with ATP) observed with lin-benzo-ATP (Table III). The $K_m$ values for lin-benzo-ATP and ATP were similar. There is no evidence for any such effect within the catalytic subunit of the cAMP-dependent protein kinase where the $V_{\text{max}}$ for lin-benzo-ATP is quite comparable to that for ATP (Hartl et al., 1983; this study) and the $K_m$ values for lin-benzo-ATP and ATP are very similar. The binding affinity of lin-benzo-ADP for the holoenzyme form (cGMP) of the cGMP-dependent enzyme is quite similar to that for the catalytic form of the enzyme (+cGMP) (Table I). By contrast, with the cAMP-dependent protein kinase, the presence of the regulatory subunit with the catalytic subunit (in the type II holoenzyme) results in a 3-fold increase in affinity of lin-benzo-ADP for the enzyme when compared with that of the catalytic subunit alone (Hartl et al., 1983).

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NUCLEOTIDE BINDING TO PROTEIN KINASE

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REFERENCES