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Contents

Contents of Number 1/2	Studies on the S-State Distribution in Euglena
Original Communications	gracilis G. H. Schmid and P. Thibault 60
A Small-Angle-Scattering Camera without Slithigh- Error, II. (In German) H. WAWRA 1 A New Red Dipyrromethene Pigment from <i>Candida</i> boidinii	Sensitive Fluorimetric Assay for Adenosine De- aminase with Formycin as Substrate; and Sub- strate and Inhibitor Properties of some Pyra- zolopyrimidine and Related Analogues J. WIERZCHOWSKI and D. SHUGAR 67
S. LANG, H. BEYER, P. SISTIG, F. WAGNER, W. TROWITZSCH, and V. WRAY 7 Ellagitannins from the Leaves of Vitis vinifera (In German) CH. KARL, G. MÜLLER, and P. ALSTED PEDERSEN 13	Elucidation of the Different Effects of Polyamines and Other Naturally-Occurring Inhibitors of Cell Proliferation (Chalones) on T-Lymphocyte and Granulocyte Colony Growth <i>in vitro</i> R. MASCHLER, C. J. SMITH, J. C. ALLEN, and H. R. MAURER 74
2-Methoxystypandrone, a New Naphthoquinone from <i>Rhamnus fallax</i> B. (In German) HW. RAUWALD and H. MIETHING 17	Effect of Organic Solvents on Enzymatic Hydro- lysis of Celluloses M. KUMAKURA and I. KAETSU 79
Enzymatic Synthesis of Sinapine from 1-O-Sinapoyl- β -D-glucose and Choline by a Cell-Free System from Developing Seeds of Red Radish (<i>Raphanus</i> sativus L. var. sativus) D. STRACK, W. KNOGGE, and B. DAHL- BENDER 21	Time-Resolved Fluorescence Studies of Hematoporphyrin in Different Solvent SystemsA. ANDREONI, R. CUBEDDU, S. DE SILVESTRI, G. JORI, P. LAPORTA, and E. REDDI83On the Origin of the Non-Haemic Iron Transferrin
Inhibition of Ergosterol Biosynthesis by Eta- conazole in Ustilago maydis E. EBERT, J. GAUDIN, W. MUECKE, K. RAM- STEINER, CH. VOGEL, and H. FUHRER 28	ESR Signal: ESR Investigations on Histidine- Iron-Ascorbic Acid Systems W. LOHMANN, D. HOLZ, B. KIEFER, and D. SCHMIDT 90 Coupling of Monomethoxypolyethyleneglycols to
Isoelectric Focusing of the NAD(P)H-Cytochrome c Reductase Subunit of Chlamydomonas reinhardii Nitrate Reductase E. FERNÁNDEZ and J. CÁRDENAS 35	Obtaming of Monomounty polyconytoing of a constraint of proteins via Active Esters E. Boccù, R. LARGAJOLLI, and F. M. VERONESE 94 OH Radical-Induced Main-Chain Scission of Poly-
The Effect of Silatranes on Growth, Photosynthetic Oxygen Evolution and Dark Respiration of <i>Cyclotella cryptica</i> (Diatomeae) (In German) R. ROTH, V. P. BARYSHOK, D. I. STOM, D. WERNER, and M. G. VORONKOV 39	 (Ribonucleic Acids) under Anoxic Conditions K. WASHINO, O. DENK, and W. SCHNABEL 100 Effect of Streptococcal Extracellular Nuclease on the Carrier Activity of RNA for Streptolysin S A. TAKETO and Y. TAKETO 107
Oxygen-Evolving Extracts from a Thermophilic Cyanobacterium Synechococcus sp. S. MIYAIRI and G. H. SCHATZ 44 Synthesis of 2'-/3'-O-Acylated Adenine Nucleotide	DNA Content and Synthesis in Several Tissues and Variation of Moulting Hormone-Level in Gryllus bimaculatus DEG (Ensifera, Insecta) F. ROMER and I. EISENBEIS 112
Analogs and Their Interactions in Photophos- phorylation G. ONUR, G. SCHÄFER, and H. STROTMANN 49	Quasielastic Light Scattering of Isolated Chro- matin (In German) В. МЕUEL and Н. Nотвонм 126

Physical and Chemical Analyses of the Mineral Substance during the Development of Two Ex- perimental Cutaneous Calcifications in Rats:	of the Stem Bark of <i>Rhamnus catharticus</i> L.: Emodin-8-O- β -gentiobioside, -glucoside and -primveroside (In German)
Topical Calciphylaxis and Topical Calcergy	HW. RAUWALD 170
H.J. TOCHON-DANGUY, G. BOIVIN, M. GEOFFROY, C. WALZER, and C. A. BAUD 135	Isolation and Structure Determination of the Pre- cursors of α - and γ -Irone and Homologous Com-
Long Term Cultures of Neural Retina and Pigment Epithelium from Newborn Rabbits	pounds from Iris pallida and Iris florentina W. KRICK, FJ. MARNER, and L. JAENICKE 179
T. TSUKAMOTO and H. LUDWIG 141	Biosynthesis of 6a-Hydroxypterocarpans: Deute- rium NMR Evidence for Direct Hydroxylation
Notes	of Pterocarpans
Ent-Kaurenoic Acid: A Diterpene as Frond Exudate on Ferns of the Genus Notholaena	S. W. BANKS and P. M. DEWICK 185
E. Wollenweber, D. Marx, J. Favre-Bonvin, and C. Brassy 146	Chemotypes of Molopospermum peloponnesiacum (L.) Koch (In German)
Identification of two Gossypetin Monomethyl	KH. KUBECZKA and I. ULLMANN 189
Ethers as Yellow Flower Pigments in the	Fungitoxic Isoflavones from Lupinus albus and other Lupinus Species
Rutaceae J. B. HARBORNE and M. BOARDLEY 148	J. L. INGHAM, S. TAHARA, and
5. D. HARBORNE AND M. DOARDLET	J. B. HARBORNE 194
Antimicrobial Activity of N,N'-bis(decylmethyl)-	
α, ω -alkanediamine Dioxides	An Improved Method of Isolation and Quantitative
F. DEVINSKY, I. LACKO, L. KRASNEC,	Determination of Vincaleucoblastine from Intact
and D. MLYNARČIK 151	Plants and Tissue Cultures of Catharanthus roseus
Fluorescence Patterns of Chromatin and Cytoplasm	G. DON. (In German) W. HOFMANN, KH. KUBECZKA, and
by Hematoxylin Solutions	FC. CZYGAN 201
J. C. STOCKERT, M. L. MOLERO,	F0. OLIGAN 201
and R. H. ESPELOSIN 153	Isolation and Identification of Sterols in the Me-
On the Construction of "Optimal" Phylogenetic	tabolism of the Fungus Botrytis cinerea
Trees	(In German)
G. DE SOETE 156	P. FLESCH, L. GWINNER, S. MÜLLER, and P. SIMON 207
	P. Simon 207
Contents of Number 3/4	Mechanisms of Inhibition by Mevinolin (MK 803) of Microsome-Bound Radish and of Partially
Original Communications	Purified Yeast HMG-CoA Reductase
Composition and Accumulation of Essential Oil in	(EC.1.1.1.34) Th. J. Bach and H. K. Lichtenthaler 212
Matricariae Radix (2. Communication) (In German)	
J. REICHLING, W. BISSON, H. BECKER,	Optically Detected Magnetic Resonance of Porphy-
and G. Schilling 159	rin Complexes in the Bacterium Rhodopseudo-
	monas sphaeroides J. BECK, J. U. VON SCHÜTZ, and H. C. WOLF 220
Flavone C-Glycoside from Metzgeria furcata	5. DECK , 5 . C . VON SCHU12 , and 11 . C . WOLF 220
(Hepaticae) (In German) R. Theodor, R. Mues, H. D. Zinsmeister,	Affinity of Phycocyanin Chromopeptides to
and K. R. Markham 165	Histidyl-Sepharose Gels: A Model for Histidine-
	Tetrapyrrol-Interactions in Biliproteins
A New Investigation on the Composition of Rhamni	J. RABIER, M. VIJAYALAKSHMI, and
cathartici Cortex, IV. The Main Anthraglycosides	W. Rüdiger 230

Sulfide Stimulation of Light-Induced Hydrogen Evolution by the Cyanobacterium Nostoc mus-		
corum H. Weisshaar and P. Böger	237	
Ribulose-1,5-bisphosphate Carboxylase-Oxygen New Aspects Respective the pH-Dependance the Carboxylation Reaction	e of	
P. BRAUN, J. BODE, and A. WILD	243	
Evidence for Unequal Misses in Oxygen Flash M Sequence in Photosynthesis MJ. DELRIEU	7 ield 247	
Calorimeter-Fermentor Combination for Investions on Microbial Steady State Cultures H. P. LEISEIFER and G. H. SCHLESER	259	
The 'Double' Michaelis-Menten Equation: Est tion of Parameters	ima-	
A. C. BORSTLAP and P. G. DOUCET	268	
L-Usnate-Urease Interactions: Binding Sites f Polymerization	or	
B. CIFUENTES, I. GARCÍA, and C. VICENTE	273	
Enhancement of <i>in-vitro</i> Translation of Eukary RNAs by Cyclic AMP	yotic	
M. E. John and W. Knöchel	277	
Characterisation of the Enzyme Intermediates of the Sarcoplasmic Transport ATPase by the Use of Inhibitors		
W. Boll and M. Makinose	282	
Thermal Stability of DNA Interacting with azolidone and Cu(II) Ions	Fur-	
M. MAITI, S. GHOSH, A. CHATTERJEE, and S. N. CHATTERJEE	290	
Circular Permutation Analysis of Phage T4 I by Electron Microscopy	ONA	
G. F. GROSSI, M. F. MACCHIATO, and G. GIALANELLA	294	
On a Quantitative Determination of Antibodic Lipids and Proteins	es to	
A. RADUNZ	297	
Effect of Sodium Cholate on the Phase Trans Temperature of Dipalmitoyl Phosphatidylch S. BANERJEE and S. N. CHATTERJEE		
On the Interaction of Phospholipid Vesicles Chaotropic Ions	with	
TH. PIGOR and R. LAWACZECK	307	

Growth Characteristics of Anaero	bically Treated
Early and Late S-Period of Ehr	lich Ascites Tu-
mor Cells after Reaeration	
R. MERZ and F. SCHNEIDER	313
Comparative Investigations on Defe and Gland Morphology of Abdo	v

and Giand Morphology of Abdommar Defensive
Glands from Rove Beetles of the Subtribe Philon
thina (Coleoptera, Staphylinidae) (In German)
K. DETTNER 319

Is the Fly Visual Pigment a Rhodopsin? K. VOGT 329

Notes

The Phytoalexin Response of Lathyrus sativu	8
(Grass Pea)	
D. J. ROBESON and J. B. HARBORNE	334
Drostanlandin Symthesis Inhibitors Deduce	Cam

Prostaglandin Synthesis Inhibitors R	Leduce Can-
nabis and Restraint Stress Induced	Increase in
Rat Brain Serotonin Concentrations	
S. K. BHATTACHARYA and	
D. BHATTACHARYA	337

Erratum to A. ANDREONI, R. CUBEDDU, S. DE SILVESTRI, G. JORI, P. LAPORTA, and E. REDDI (Z. Naturforsch. 38c, 83-89 (1983) 338

Contents of Number 5/6

Original Communications

The Stereochemistry of Silybin (In German)	
H. LOTTER and H. WAGNER	339

- Identification of Anthocyanins and Intermediates of Anthocyanin Biosynthesis from *Petunia hybrida* Using High Performance Liquid Chromatography A. W. SCHRAM, L. M. V. JONSSON, and P. DE VLAMING 342
- Formation of Benzophenanthridine Alkaloids by Suspension Cultures of *Eschscholtzia californica* J. BERLIN, E. FORCHE, V. WRAY, J. HAMMER, and W. HÖSEL 346
- Circulardichroism of C-Phycocyanin: Origin of Optical Activity in Denatured Biliproteins and Evidence for an Intermediate during Unfolding H. LEHNER and H. SCHEER 353

Contents

Chromopeptides from Phytochrome and Phyco- cyanin. NMR Studies of the P_{fr} and P_r Chromo- phore of Phytochrome and E , Z Isomeric Chro- mophores of Phycocyanin F. THÜMMLER, W. RÜDIGER, E. CMIEL, and S. SCHNEIDER 359	Characteristic Features of the Regulatory Functions of the D-Ribulose 1,5-Bisphosphate Carboxylase/ Oxygenase from Spinach J. VATER, T. GAUDSZUN, B. LANGE, N. ERDIN, and J. SALNIKOW 418
Prunasin Biosynthesis by Cell-Free Extracts from Black Cherry (<i>Prunus serotina</i> Ehrh.) Fruits and Leaves	A Cysteine Desulfhydrase Specific for D-Cysteine from the Green Alga <i>Chlorella fusca</i> A. SCHMIDT and I. ERDLE 428
J. E. POULTON and SUN-IN SHIN 369	Regulation of Nitrogenase Activity in <i>Rhodopseudo</i> -
Comparison of Purine Metabolism in Suspension Cultured Cells of Different Growth Phases and	monas capsulata AD 2 K. ALEF 436
Stem Tissue of Catharanthus roseusF. HIROSE and H. ASHIHARA375	Nitrate Reductase from a Mutant Strain of Chlamy- domonas reinhardii Incapable of Nitrate Assimila-
Uptake and Conversion of Three Different 5-Ring Polycyclic Aromatic Hydrocarbons (PAHs) in Call Summarian Cultures of Various Changes	tion E. Fernández and J. Cárdenas 439
Cell Suspension Cultures of Various Chenopo- diaceae-Species H. HARMS 382	A Membrane Bound Cysteine Oxydase from the Cyanobacterium Synechococcus 6301
Geranyl Derivatives as Inhibitors of the Caroteno-	A. SCHMIDT and E. KRÄMER 446
genesis in Synechococcus PCC 6911 (Cyanobacteria) F. JÜTTNER and O. BOGENSCHÜTZ 387	A Specific and Sensitive Radioimmunoassay for Human Choriogonadotropin
Interconversion of Carotenoids and Quinones after Onset of Photosynthesis in Chloroplasts of Higher Plants	K. MURALIDHAR, G. CHAUDHURI, J. LIPPES, and O. P. BAHL 451
K. H. GRUMBACH 393	Effect of Pressure on the Absorption Spectra of Phycobiliprotein and Porphyridium cruentum
The Influence of the Glycerol-3-Phosphate Level in the Stroma Space on Lipid Synthesis of Intact	S. S. BRODY and L. STELZIG 458
Chloroplasts A. SAUER and KP. HEISE 399	Bioalkylation of Nucleic Acids in Mice by Insecti- cides, I. Alkylation of Liver RNA and DNA by Chlorpyrifos
Inability of Petite Mutants of Industrial Yeasts to	I. Y. MOSTAFA, Y. M. ADAM, and
Utilize Various Sugars, and a Comparison with the Ability of the Parent Strains to Ferment the	S. M. A. D. ZAYED 461
the Same Sugars Microaerophilically J. F. T. SPENCER, D. M. SPENCER, and R. MILLER 405	Electromagnetic Modulation of Biological Processes: Bicarbonate Effect and Mechanistic Considera- tions in the Ca-Uptake by Embryonal Chick
Studies on the Regulatory Role of <i>trans</i> -Cinnamic Acid on the Activity of the Phenylalanine Am-	Tibia in vitroG. COLACICCO and A. A. PILLA465
monia-Lyase (PAL) in Suspension Cultures ofDaucus carota L.W. Noé and H. U. SEITZ408	Electromagnetic Modulation of Biological Processes: ATPase Function and DNA Production by Raji Cancer Cells <i>in vitro</i>
Meta-Methylation of Flavonol Rings $A(8-)$ and $B(3'-)$ Is Catalysed by Two Distinct O-Methyl-	G. COLACICCO and A. A. PILLA 468
transferases in <i>Lotus corniculatus</i> M. JAY, V. DE LUCA, and R. IBRAHIM 413	The Influence of the Extracellular Concentration of Calcium, Magnesium and Sodium on the Repo-

larizing Phase of the I	Receptor	Potential	of	the
Limulus Ventral Nerve	Photore	eceptor		
H. STIEVE, M. BRUNS,	and H	. Gaube		471

Notes

Adsorbent Culture – Method for the 1	\mathbf{Enh}	anced
Production of Secondary Substances	in	Plant
Suspension Cultures (In German)		
B. KNOOP and R. BEIDERBECK		484

- The Structure of a Ring-A-seco-nor-Triterpenol from Hoya australis Leaf Wax W. J. BAAS 487
- Studies on the Distribution of Phytosterols in Dioscorea Species Tubers P. G. KADKADE, C. LUJAN, and C. ROLZ 488
- Quercetin-3-α-L-(2-O-α-L-rhamnopyranosidoarabopyranoside), a New Flavonolglycoside from Seeds of *Brassica nigra* (L.) Koch. (In German) H. GEIGER, H. MAIER, and K. R. MARKHAM 490
- Verpacrocin, a Polyene Pigment from Mycelial Cultures of Verpa digitaliformis (Pers.) Fr. (Ascomycetes) (In German)
 H. BESL, A. BRESINSKY, B. MEIXNER,
 U. MOCEK, and W. STEGLICH 492
- Inheritance of Sesquiterpenoid Phenolic Acid
 Esters (Guayulins) in F₁ Hybrids of Parthenium (Asteraceae)
 H. M. BEHL, B. MARCHAND, and
- E. RODRIGUEZ 494 Chemical Composition of the Secretion from the Anal Sacs of *Civettictis civetta* (Schreber, 1776)
- J. JACOB and H. SCHLIEMANN 497 The Possible Common Origin of tRNA and 5S rRNA

M. UJHELYI

Rotation of Cells and Ion Exchange Beads in the MHz-Frequency Range G. KÜPPERS, B. WENDT, and U. ZIMMERMANN 505

501

Temperature Regulation in Wood Ant Nests (Formica polyctena Förster) (In German) K. HORSTMANN 508

Contents of Number 7/8

Original Communications

- Crystal Packing of Odd-Chain Saturated Triglycerides D. L. DORSET 511
- New Cembranoids from Tobacco, II A.-M. Bylov, U. Brümmer, W. Hass, F. See-HOFER, V. HEEMANN, and V. SINNWELL 515
- 3,7,11,15-Cembratetraen-6-ol, a New Cembranoid from Tobacco, III V. HEEMANN, A.-M. BYLOV, U. BRÜMMER, W. HASS, and F. SEEHOFER 517
- Synthesis of Mono-, Di- and Trigalloyl- β -D-glucose by β -Glucogallin-Dependent Galloyltransferases from Oak Leaves G. G. GROSS 519
- Carboxylic Acids in Wheat, Rye and Barley E. LOHAUS, I. BLOS, and W. RÜDIGER 524
- Separation, Purification, and Characterization of Two Isoforms of Glutamine Synthetase from *Chlamydomonas reinhardii*F. J. FLORENCIO and J. M. VEGA 531
- Transformations of Phenazines in *Pseudomonas* aureofaciens (In German) A. RÖMER and E. LANGE 539

Differential Regulation of Flavone Glycosylation during Ontogeny of *Silene pratensis* J. M. STEYNS, G. VAN NIGTEVECHT, G. J. NIE-MANN, and J. V. BREDERODE 544

- Ontogeny and Biosynthesis of Isovitexin 7-O-Galactoside in a Mutant of *Silene pratensis* Unable to Glycosylate this Compound in the Petals J. VAN BREDERODE and J. M. STEYNS 549
- Relationship between Flower Development, Anthocyanin Accumulation and Activity of Enzymes Involved in Flavonoid Biosynthesis in Matthiola incana R. Br.
 B. DANGELMAYR, G. STOTZ, R. SPRIBILLE,
 - and G. FORKMANN 551

556

Variable Fluorescence and Fluorescence Spectra of Algae after Herbicide-Induced Pigment Bleaching

N. V. KARAPETYAN, R. STRASSER, and P. Böger

- Investigations on the Synthesis and Biological Significance of Glutamic-1-semialdehyde as a Precursor of the Chlorophylls (In German) H.-U. MEISCH and R. MAUS 563
- Nitrogen Fixation and Nitrate Respiration by Azospirillum brasilense H. BOTHE, G. BARBOSA, and J. DÖBEREINER 571
- Cyclic AMP Is Not Involved as a Second Messenger in the Response of Soybean to Infection by Phytophthora megasperma f. sp. glycinea M. G. HAHN and H. GRISEBACH 578
- Biosynthesis of Cyanogenic Glycosides in Butterflies and Moths: Incorporation of Valine and Isoleucine into Linamarin and Lotaustralin by Zygaena and Heliconius Species (Lepidoptera) V. WRAY, R. H. DAVIS, and A. NAHRSTEDT 583
- Caffeine-Induced Surface Blebbing and Budding in the Acellular Slime Mold Physarum polycephalum J. KUKULIES, W. STOCKEM, and K. E. WOHLFARTH-BOTTERMANN 589
- Regulation of Light Energy Distribution between Photosynthetic Pigment Systems; a Possible Role of Leaf Anatomy G. HARNISCHFEGER and G. ZENK 600
- Energy Parameters, Macromolecular Synthesis and Cell Cycle Progression of in vitro Grown Ehrlich Ascites Tumor Cells after Inhibition of Oxydative ATP Synthesis by Oligomycin W. KROLL, M. LÖFFLER, and F. SCHNEIDER 604
- Embryonic Hemoglobins in Mammals: The Primary Structures of ζ -, ε - and ϑ -Chains of the Pig (Sus scrota domestica) (In German) F. A. BIEBER, H. ASCHAUER, S. M. BEKTAS, and G. BRAUNITZER 613

- Studies on the Transport of Anions and Zwitterions of Acidic Amino Acids in Streptomyces hydrogenans J. FRITSCH and W. GROSS 617
- Hydrogen Bond Indices and Tertiary Structure of Yeast tRNA^{Phe} M. S. DE GIAMBIAGI, M. GIAMBIAGI, and D. M. S. ESQUIVEL 621
- Effect of 5'-Terminated (2'-5')-Oligoadenylates on Cap Degrading Activities in Rat Liver Nuclei W. MICHELS and E. SCHLIMME 631
- Decreased Production of RNA-Streptolysin S in Streptococci Devoid of Extracellular RNase Activity
 - A. TAKETO and Y. TAKETO 635
- Activation and Inhibition of Na/K-ATPase by Filipin-Cholesterol Complexation. A Correlative Biochemical and Ultrastructural Study on the Microsomal and Purified Enzyme of the Avian Salt Gland D. GASSNER and H. KOMNICK 640
- Incorporation of a Voltage Sensitive Pore from Guinea Pig Heart Mitochondria into Black Lipid Membranes and Characterization of Electrical Properties W. SCHREIBMAYER, H. HAGAUER, and 664
 - H. A. TRITTHART

Notes

- Flavonoids from the Leaf Resin of Hymenoclea salsola T.&G. (Asteraceae) P. PROKSCH, E. WOLLENWEBER, and E. RODRI-668 GUEZ
- New Papaveraceae Alkaloids Including an 7,8-Oxygenated Isoquinoline, the Postulated Precursor of the Cularine Alkaloids G. BLASCHKE and G. SCRIBA 670
- Conical Kinetochores with Single Microtubules in the Micronuclear Mitosis of the Ciliate Colpoda steinii
 - 672 A. KUCK and A. RUTHMANN
- Time-Shifted Mating Periods in Two Closely Related Corixid Species 675 J. THEISS

Contents of Number 9/10	Effects on Pigment Accumulation and Fluores- cence Properties
Original Communications	G. LASKAY, E. LEHOCZKI, I. MARÓTI, and L. SZA-
Epicuticular Waxes from Leaves and Stems of Jojoba (Simmondsia chinensis [Link] Schneider) PG. GÜLZ and C. EICH 679	LAY 736 Effects of Pyridazinone Herbicides during Chloro- plast Development in Detached Barley Leaves. II.
Chemistry and Morphology of Epicuticular Waxes from Various Organs of Jojoba (Simmondsia chi- nensis [Link] Schneider) PG. GÜLZ and K. HANGST 683	Effects on Lipid Content, Fatty Acid Composition and Ultrastructure of Chloroplasts G. LASKAY, T. FARKAS, E. LEHOCZKI, and K. GULYA 741
Isolation and Structural Determination of a New Methylated Triterpenoid from Rhizomes of <i>Iris</i> versicolor L.	Chemistry of Bacteriochlorophyll b: Identification of Some (Photo)Oxidation Products R. STEINER, E. CMIEL, and H. SCHEER 748
W. KRICK, FJ. MARNER, and L. JAENICKE 689	Urobiliverdin, a New Bile Pigment Deriving from
Antifungal Phytoalexins in <i>Phaseolus aureus</i> Roxb. M. J. O'NEILL, S. A. ADESANYA, and M. F. Ro-	Uroporphyrin E. BENEDIKT and HP. KÖST 753
BERTS 693	Conformational Study of Catecholamines in Solu-
Aureol and Phaseol, Two New Coumestans from <i>Phaseolus aureus</i> Rosb. M. J. O'NEILL 698	tion P. Šolmajer, D. Kocjan, and T. Šolmajer 758
Synthesis and Biological Evaluation of 7a,7a,7a,8a,8a,8a-Hexafluororiboflavin and 7a,7a,7a,8a,8a,8a-Hexafluoro-FMN	The Effects of Tumor Sera on Cell Shape and Photo- synthesis of Euglena gracilisH. G. RUPPEL and B. BENNINGHOFF763
P. Nielsen, A. Bacher, D. Darling, and M. Cushman 701	Lipids and Enzymatic Activities in Vacuolar Mem- branes Isolated via Protoplasts from Oat Primary
3'-Hydroxylation of 4'-Methoxyisoflavones by Fusa- rium oxysporum f. lycopersici K. MACKENBROCK and W. BARZ 708	Leaves B. Verhoek, R. Haas, K. Wrage, M. Linscheid, and E. Heinz 770
Kinetin-Mediated Stimulation of Accumulation of Buckwheat Flavonoids in the Dark U. MARGNA and T. VAINJÄRV 711	A Study on Oxygen Evolution and on the S-State Distribution in Thylakoid Preparations of the Fil- amentous Blue-Green Alga Oscillatoria chalybea K. P. BADER, P. THIBAULT, and G. H. SCHMID 778
Uptake and Release of Abscisic Acid by RunnerBean Root Tip SegmentsW. HARTUNG and B. DIERICH719	The Use of O ₂ -Evolving Subchloroplast Particles to Study Acceptor and Inhibitor Sites on the Re-
Distribution and Variation of the Catalytic Activity of the Enzyme Responsible for Carthamin Synthe- sis in Safflower Seedlings	ducing Side of Photosystem II W. S. COHEN and J. R. BARTON 793
K. SAITO, Y. TAKAHASHI, and M. WADA 724	Electroimmunodiffusion – a Powerful Tool for Quantitative Determinations of Both Soluble and
Further Studies on Induction of Enzymes of Phyto- alexin Synthesis in Soybean and Cultured Soy- bean Cells	Membrane Bound Chloroplast ATPase, CF ₁ P. Roos and R. J. BERZBORN 799
J. LEUBE and H. GRISEBACH 730	Analogue of Hippuran: Labelled Ruthenocenoyl- glycin. Synthesis, Organ Distribution and Clear-
Effects of Pyridazinone Herbicides during Chloro- plast Development in Detached Barley Leaves. I.	ance (In German) M. WENZEL und G. SCHACHSCHNEIDER 806

Contents

Immobilization of <i>E. coli</i> Cell as an Antigen by Ra- diation Polymerization Method	F
I. KAETSU, M. KUMAKURA, S. KIKUCHI, S. ADACHI, and M. Suzuki 812	
Exchange of the Structural Water of Collagen and Alcanol-Induced Molecular Rearrangement (In German)	S
TH. NEMETSCHEK, E. KNÖRZER, W. FOLKHARD, W. GEERKEN, K. JELINEK, C. KUHLEMANN, E. MOs- LER, and H. NEMETSCHEK-GANSLER 815	F
Preparation and Characterization of Actin from Liver	E
B. Jaberg 829	
Transient Activation of the Ca ²⁺ -ATPase from Sar- coplasmic Reticulum during Thiol Modification by 5,5'-Dithiobis(2-nitrobenzoate)	A
G. SWOBODA and W. HASSELBACH 834	
Phosphorylation of Ca-ATPase of Sarcoplasmic Re- ticulum with Different Substrates	A
H. BARRABIN and L. DE MEIS 845	
 Composition of the Heterogeneous Sex Attracting Secretion of the Dung Beetle, <i>Kheper lamarcki</i> B. V. BURGER, Z. MUNRO, M. RÖTH, H. S. C. SPIES, V. TRUTER, G. D. TRIBE, and R. M. CREWE 848 	ſ
Stimulation of Juvenile Hormone Biosynthesis in vitro by Locust Allatotropin	F
HJ. FERENZ and I. DIEHL 856	
Notes	
Interaction of Ascorbic Acid with Disulfides	
J. E. Fleming, K. G. Bensch, J. Schreiber, and W. Lohmann 859	C
ESR Investigations on the Modification of the Ascor- bic Acid-Erythrocyte Interaction by Ferrous Ions W. LOHMANN, B. KIEFER, D. HOLZ, D. SCHMIDT,	5
and J. SCHREIBER 862	

d
L.
4

Photosensitized Effects of Furocoumarins: the Pos- sible Role of Singlet Oxygen D. VEDALDI, F. DALL'ACQUA, A. GENNARO, and G. RODIGHIERO 866
Supplementary Data on the Sex Attractant System of Panolis flammea E. PRIESNER and M. SCHROTH 870
Receptors for Di-Unsaturated Pheromone Analo- gues in the Male Summerfruit Tortrix Moth E. PRIESNER 874
Effect of Ecdysterone on Food Intake of Locusta mi- gratoria Hoppers P. J. RAO and H. REMBOLD 878
Aminoacyl-tRNA Synthetases in Liver, Spleen and Small Intestine of Aged Leukemic and Aged Nor- mal Mice HJ. GABIUS, S. GABIUS, G. GRAUPNER, F. CRA- MER, and S. REHM 881
A Non-Thermal Effect of Millimeter Wave Radiation on the Puffing of Giant Chromosomes C. Koschnitzke, F. Kremer, L. Santo, P. Quick and A. Poglitsch 883
Determination of Measles Virus Protein Molecular Weights on High Percentage, Highly Cross-Linked SDS Polyacrylamide Gels M. J. CARTER and K. BACZKO 887
Flavonoids from the Leaf Resin of Hymenoclea sal- sola T. & G. (Asteraceae) P. PROKSCH, E. WOLLENWEBER, and E. RODRI- GUEZ 668
Contents of Number 11/12
Original Communications
5,7,3',4',5'-Pentahydroxyflavanone in the Bracts of Helichrysum bracteatum G. FORKMANN 891

The Flavones of the European Species of *Silene* Section Elisanthe

O. MASTENBROEK, J. J. KNORR, R. KAMPS-HEINS-BROEK, J. W. MAAS, J. M. STEYNS, and J. VAN BRE-DERODE 894

Х

Isolation and Identification of Isoflavanone Phyto- alexins from Leaflets of <i>Diphysa robinioides</i>	Cyt t
J. L. INGHAM and S. TAHARA 899	<i>g</i> U
Wounding-Induced Increase of Quinolizidine Alka- loid Accumulation in Lupin Leaves	Мо
М. Wink 905	F L
Site of Action of Growth Inhibitory Tryptophan Analogues in <i>Catharanthus roseus</i> Cell Suspen- sion Cultures	The A
F. SASSE, M. BUCHHOLZ, and J. BERLIN 910	t F
Selection of Cell Lines of Catharanthus roseus with Increased Tryptophan Decarboxylase Activity F. SASSE, M. BUCHHOLZ, and J. BERLIN916	The
 Structure of Ascorbic Acid and Its Biological Function: V. Transport of Ascorbate and Isoascorbate across Artificial Membranes as Studied by the Spin Label Technique W. LOHMANN and J. WINZENBURG 923 	H The C
On the Origin of the Non-Haemic Iron Transferrin ESR Signal: ESR Investigations on Histidine-Iron- Citric Acid Systems W. LOHMANN, D. HOLZ, B. KIEFER, and D. SCHMIDT 926	N S
Enhancement Effect of Hydrophobic Vinyl Com- pounds in Enzymatic Hydrolysis of Waste Papers M. KUMAKURA and I. KAETSU 929	a
Purification and Characterization of a Dissimilatory Nitrite Reductase from the Phototrophic Bacteri- um <i>Rhodopseudomonas palustris</i> M. PREUSS and JH. KLEMME 933	Dis r
Isolation and Fast Purification of Neocarzinostatin by FPLC-Ion Exchange Chromatography D. DENKLAU, W. KÖHNLEIN, G. LÜDERS, and J. STELLMACH 939	I
 ¹⁵N-Coprobiliverdin, a New Model Chromophore HP. Köst, E. BENEDIKT, E. CMIEL, and S. SCHNEIDER 943 	
Circular Dichroism of Chromopeptides from Phyco- cyanin	
C. Scharnagl, E. Köst-Reyes, S. Schneider, H P. Köst, and H. Scheer 951	

Cytochromes and Anaerobic Sulfide Oxidation in the Purple Sulfur Bacterium Chromatium warmin- gii
U. WERMTER and U. FISCHER 960
Molecular Properties of High Potential Iron Sulfur Protein of Chromatium warmingii
U. WERMTER and U. FISCHER 968
The Phycobiliproteids in <i>Cyanophora paradoxa</i> as Accessoric Pigments and Nitrogen Storage Pro- teins
H. E. A. SCHENK, J. HANF, and M. NEU-MÜLLER 972
The Influence of Cycloheximide and Chlorampheni- col on the Biosynthesis of the Photosynthetic Pig- ments in Cyanophora paradoxa. I. Photosynthetic Oxygen Evolution (In German)
H. E. A. SCHENK and M. NEU-MÜLLER 978
The Influence of Cycloheximide and Chlorampheni- col on the Biosynthesis of the Photosynthetic Pig- ments in <i>Cyanophora paradoxa</i> . II. Carotenoids (In German)
M. NEU-MÜLLER, H. E. A. SCHENK, and H. STRANSKY 984
Che Influence of Cycloheximide and Chlorampheni- col on the Biosynthesis of the Photosynthetic Pig- ments in Cyanophora paradoxa. III. Chlorophyll a and Phycochromoproteids (In German) M. NEU-MÜLLER and H. E. A. SCHENK 990
Distribution of Chlorophylls, Carotenoids and Qui- nones in Chloroplasts of Higher Plants K. H. GRUMBACH 996
Chlorophyll-Lipid-Interactions in Monomolecular Layers H. HEITHIER and H. MÖHWALD 1003
Sex Pheromone Components of the Gamma Moth,
Autographa gamma (L.) (Lepidoptera: Noctuidae)E. DUNKELBLUM and S. GOTHILF1011
A Conformational Transition of the Sarcoplasmic Reticulum Calcium Transport ATPase Induced by Vanadate
W. HASSELBACH, P. MEDDA, A. MIGALA, and B. Agostini 1015

Contents

Enzymatic Removal of O ⁶ -Ethylguanine versus Sta- bility of O ⁴ -Ethylthymine in the DNA of Rat Tis-	Incorporation of ¹⁵ N-Ammonia into Serotonin in Cotyledons of Maturing Walnuts
sues Exposed to the Carcinogen Ethylnitrosourea: Possible Interference of Guanine-O ⁶ Alkylation	W. GROSSE and F. ARTIGAS 1057
with 5-Cytosine Methylation in the DNA of Repli- cating Target Cells	N-Terminal Sequence of a Porphobilinogen-Synthase (In German)
R. MÜLLER and M. F. RAJEWSKY 1023	B. LINGNER and T. KLEINSCHMIDT 1059
Rate of OH Radical Induced Strand Break Forma- tion in Single Stranded DNA under Anoxic Con- ditions. An Investigation in Aqueous Solutions	Sequence Analysis of the Cloned Cucumis melo Highly Repetitive Satellite DNA A. BRENNICKE and V. HEMLEBEN 1062
Using Conductivity Methods	A. DRENNICKE allu V. HEMLEBEN 1002
E. Bothe, G. A. Qureshi, and D. Schulte-Froh- LINDE 1030	Variation in Cephalic Volatile Substances in Rela- tion to Worker Age and Behavior in the Stingless Bee, Scaptotrigona postica
The Intensity Dependence of the Receptor Potential of the <i>Limulus</i> Ventral Nerve Photoreceptor in Two Defined States of Light- and Dark Adapta-	W. FRANCKE, W. SCHRÖDER, E. ENGELS, and W. ENGELS 1066
tion H. Stieve, M. Bruns, and H. Gaube 1043	Antibodies against the α -Factor Pheromone of Sac- charomyces cerevisia
	U. TILLMANN and H. HAHN 1069
Notes	A New Cultural Method for <i>Trichoplax adhaerens</i> F. E. SCHULZE (In German)
Identification of the Major Anthocyanin of Carrot Cells in Tissue Culture as Cynidin 3-(Sinapoylxy-	K. G. Grell 1072
losylglucosylgalactoside) J. B. Harborne, A. M. Mayer, and N. Bar-Nun	Subject Index 1073
1055	Authors Index 1095

XII

Circulardichroism of C-Phycocyanin: Origin of Optical Activity in Denatured Biliproteins and Evidence for an Intermediate during Unfolding

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Phycocyanin, Phycoerythrin, Phytochrom, Circulardichroism, Denaturation

The circulardichroism spectra (cd) of native and (partially) urea-denaturated C-phycocyanin (PC) in the spectral range 700-210 nm are presented. The large ellipticities observed in the chromophoric region of native PC are retained in the fully denatured state (8 M urea). This is similar to the behavior of the red form of phytochrome (P_r), but in contrast to C-phycocrythrin (PE). These differences are rationalized in terms of epimeric equilibria between P- and M-helix shaped chromophores. Depending on the number and location of the chirality centers present in the tetrapyrrol moities *a priori*, the excess populations of the inherently chiral P- and M-helices differ, thus accounting for large (PC, P_r) or small (PE) ellipticities in the denatured pigments. Hence, the large optical activity observed for the former is generated by an excess population of the P-helix induced by the asymmetric C-2, C-3 and C-3'. In PE the additional chirality center at C-16 counteracts the influence of the others. The excess population of the enarry vanishing cd reported for denatured PE.

The cd has also been studied at intermediate urea concentrations. Unfolding of PC with urea can be interpreted from these data as a stepwise process. Monitoring the urea induced unfolding of PC by cd at different wavelengths ($\lambda = 220$, 345, and 610 nm); the "melting point" of the apoprotein (4-5 M urea) coincides with the extrema of the titration curves obtained in the chromophore region (345, 610 nm). These results give direct evidence for the existence of an intermediate species whose population reaches a maximum at 4-5 M urea.

Introduction

Phycobiliproteins are light harvesting pigments of certain algae (for some recent reviews, see *e.g.* refs. 1-4).

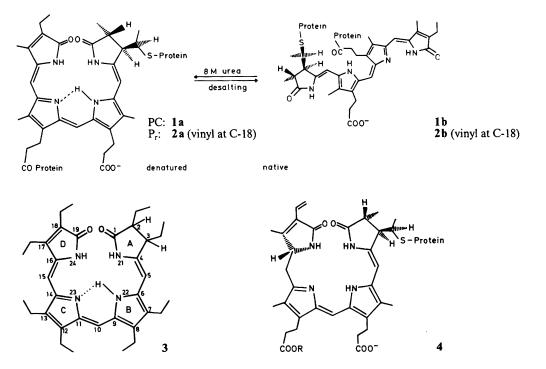
They contain chromophores like 1 covalently bound to the apoproteins. Whereas the denatured pigments have properties typical to free bile pigments of similar structure, *e.g.* **3**, these properties are conspicously changed in the native pigments, in an adaptation to their function. From UV-Vis studies and complementary MO-calculations it has been concluded, that these changes are in part due to a conformational transition from a cyclic helical (*e.g.* **1a**) to a more extended conformation (*e.g.* **1b**,

Reprint requests to Prof. Dr. H. Scheer. 0341-0382/83/0500-0353 \$ 01.30/0 for leading references, see refs. 4), and that this process involves at least one intermediate state [5]. The denaturation of PE [6] and phytochrome P_r and P_{fr} [7] has been studied in detail by a combinatory evaluation of absorption, fluorescence, and circulardichroism spectroscopy. It could be shown that these techniques are complementary in exhibiting different sensitivities to the processes involved during denaturation. As an example in PE both the fluorescence and the cd-signals of the chromophores [4] decrease to almost zero upon complete unfolding of the protein [6]. On the other hand, the optical activity of phytochrome (P_r , 2) is retained upon unfolding of the protein [7].

We here wish to report the corresponding cd spectra for the urea-induced denaturation of PC. The results obtained permit conclusions with respect to the origin of optical activity of biliproteins in general, especially in their denaturated states. Moreover, more direct evidence has been obtained for a stepwise unfolding of PC proposed earlier from UV-Vis data [5].

Abbreviation: PC, C-phycocyanin; PE, C-phycoerythrin; Pr, red absorbing form of phytochrome; cd, circulardichroism; UV-Vis, absorption spectroscopy in the ultraviolett and visible range.

H. Lehner and H. Scheer · Circulardichroism of C-Phycocyanin



Experimental part

PC from Spirulina platensis was prepared by the method reported earlier [8]. Urea was reagent grade (Merck, Darmstadt) or ultrapure (Schwarz & Mann, New York). In the spectroscopic studies, stock solutions of PC ($\approx 20 \,\mu$ M) in potassium phosphate buffer (0.05 M, pH 7) were mixed with this buffer, and with buffer containing urea (8 M) in the appropriate ratios. UV-Vis and fluorescence spectra were recorded on a model DMR22 (Zeiss, Oberkochen) spectrometer equipped with thermostated cell holder, at 20 °C. Cd-spectra were recorded on a dichrograph Mark III (Jobin-Yvon, France) using thermostated (20 \pm 1 °C) cylindrical quartz cuvettes with variable path length (10-0.1 mm). The base lines were obtained by measuring the appropriate buffer-urea solutions under the same conditions.

Results

The cd of native PC (Fig. 1) has a positive longwavelength band peaking at 632 nm and a negative near-uv cd-band peaking at 342 nm, which are due to the chromophores. The bands below 280 nm arise primarily from the apoprotein. Both chromophore bands are structured in a similar fashion as the absorption bands (Fig. 1), although there are small but distinct variations between the cd- and UV-Vis maxima (*vide infra*).

PC contains three chromophores of identical molecular structure (1), but in different protein environments (see refs. 1-4). The fine structure of

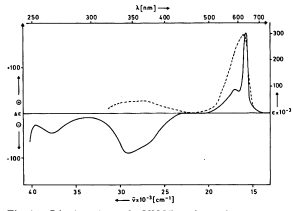


Fig. 1. Cd (----) and UV-Vis absorption spectrum (-----) of PC (4×10^{-6} M) in the chromophoric region in phosphate buffer (0.05 M, pH 7) at room temperature. Molar extinction coefficients (ϵ) and $\Delta\epsilon$ -values are given for the monomer containing three chromophores.



H. Lehner and H. Scheer · Circulardichroism of C-Phycocyanin

the long-wavelength cd-band is roughly similar to that of the absorption band. In the latter, the major band corresponds to two chromophores, one each on the α - and β -chain. The shoulder at shorter wavelengths which is resolve as a second band at low temperatures [9] arises from the second chromophore on the β -chain. This assignment is likely from the spectra of the isolated subunits (Scheer *et al.*, unpublished). The similarity of the cd-band shape to this pattern indicates a similar differentiation. Upon denaturation, the long-wavelength cd-band ($\lambda_{max} = 615$ nm) exerts a hypsochromic shift by

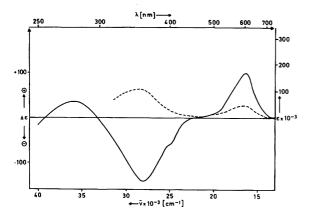


Fig. 2. Cd (—) and UV-Vis absorption spectrum (-----) of PC $(4 \times 10^{-6} \text{ M})$ in phosphate buffer (0.05 M, pH 7) containing urea (8 M) at room temperature. See Fig. 1 for molar extinction coefficients and $\Delta \varepsilon$ -values.

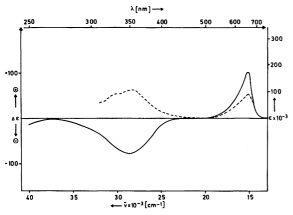


Fig. 3. Cd (----) and UV-Vis absorption spectrum (-----) of PC $(4 \times 10^{-6} \text{ M})$ in aqueous urea (8 M) titrated with HCl to pH 1.5, at room temperature. See Fig. 1 for molar extinction coefficients and $\Delta \varepsilon$ -values.

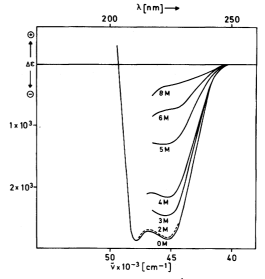


Fig. 4. Cd spectra of PC $(4 \times 10^{-6} \text{ M})$ in the peptide absorption region at various urea concentrations (0 M, 2 M, 3 M, 4 M, 5 M, 6 M, and 8 M) in 0.05 M phosphate buffer at pH 7; room temperature.

17 nm, and the $\Delta \varepsilon$ -value is decreased only by about 40% (Fig. 2). The changes in the negative band at 342 nm are less pronounced, there is even a slight increase of the ellipticities. Both bands loose their fine structure with increasing urea concentrations.

For a comparison, the cd of PC in 8 M urea at pH 1.5 is given in Fig. 3. The chromophhore is present in its protonated form under these conditions. In the UV-Vis spectrum, this leads to an increased absorption and a concomitant red-shift of the long-wavelength band [10], whereas the near-uv band remains almost unchanged. The red-shift is matched by the cd-band. The anisotropy factor of both bands is decreased by a factor of two.

The region of the peptide cd-bands (200-250 nm)shows a steady decrease with increasing urea concentrations (Figs. 4, 5), with an inflection at about 4.5 M urea. The cd-titration curves at wavelengths $\ge 300 \text{ nm}$ (corresponding to chromophore absorptions) do not show similarly steady changes but rather irregularities which are most pronounced in the region between 4 and 5 M urea. Two such curves are shown in Fig. 5, and phenomenologically similar curves are obtained at any other wavelength corresponding to chromophore rather than peptide absorptions.

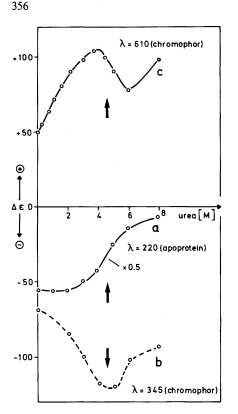


Fig. 5. Titration curves of PC with urea at three wavelengths: a: 220, b: 345 and c: 610 nm, monitored by cd.

Discussion

a. Noncoincidences between cdand UV-Vis absorption extrema

Whereas the cd fine structure in the native pigment can be interpreted like the electron absorption spectra by the three chromophores present in different environments (see refs. 1-4), the bathochromic shift of the cd-spectra with respect to the electron absorption spectra in the chromophoric region needs further discussion.

Principally, it may arise from several origins: i) A dynamic equilibrium between different conformers of each chromophore, with the populations absorbing more to the red exhibiting a stronger optical activity. ii) Each of the three chromophores is in a rather distinct environment, but the one absorbing mostly to the red has a proportionally low absorption and high optical activity. This situation is similar to (i), but represents a frozen rather than a

H. Lehner and H. Scheer · Circulardichroism of C-Phycocyanin

dynamic state. iii) At least two of the chromophores are close enough and appropriately oriented to each other to produce an exciton couplet with a positive band at the long-wavelength side.

Whereas several lines of evidence (see ref. 4) render the possibility (i) unlikely, both (ii) and (iii) are compatible with results assigning each chromophore type to a rather distinct and rigid site. Exciton coupling (e.g. iii) seems at least possible from the following arguments: In native PC the long-wavelength cd-maximum occurs at 632 nm, which is shifted by 306 cm^{-1} (12 nm) to the red as compared to the long-wavelength absorption maximum. A corresponding negative extremum would be expected around 608 nm. The pronounced dip in the cd-curve at 602 nm is indicative of this and may arise from a superposition of the negative part of the couplet with the positive band due to the "s"-chromophore. The opposite changes around 630 (negative) and 600 nm (positive) upon denaturation would also support this explanation. Excition splittings have earlier been proposed for a cryptophytan PC [11] and several allophycocyanins [12], and also discussed for PE [6], but only in the former case there exists more than circumstantial evidence.

b) Origin of optical activity in native and denatured biliproteins

The spectral properties and the conformation of the prosthetic groups of native biliproteins are strongly influenced by the corresponding apoproteins to which they are covalently bound. Thus, the chromophores exhibit an extended conformation which is chiral and rigid due to non covalent interactions with the apoprotein. On destroying the influence of the apoprotein by denaturation with urea, the conformation of the tetrapyrrol chromophores changes to a cyclic, helical arrangement (see ref. 4 for leading references). Hence, denatured biliprotein-chromophores predominantly adopt a cyclic helical conformation (2 a and 1 a respectively). In open-chain tetrapyrrols lacking any chiral influence the P- and M-helices are enantiomeric and thus equipopulated. Since the inversion barrier of the tetrapyrrol-helix is low $(42 \text{ kJ} \cdot \text{mol}^{-1} \text{ in biliver-}$ din [13]) and inversion fast at ambient temperature. the P/M-helix ratio can be changed by exerting a diastereoisomeric relationship on it. Since such

H. Lehner and H. Scheer · Circulardichroism of C-Phycocyanin

helices represent inherently chiral chromophores large optical activity is to be expected even if the excess population is small. An example for an external chiral influence represents the cd of achiral bile pigments complexed with the protein, serum albumin [14] or dissolved in chiral, optically active solvents [15-17].

In the denatured state of biliproteins the optical activity observed, is governed mainly by internal chiral influences, viz. of the chirality centers present in the chromophores. The PC chromophores possess chirality centers at C-2, C-3 and C-31 inducing a twist of the C-5 methine bridge, which is energetically favoured in one distinct direction. Hence, the large optical activity of even denatured PC may be rationalized in terms of an excess population of the P-helical arrangement being diastereoisomeric to the less populated M-helix. This interpretation is further corroborated by recent Ramachandran-type calculations [18, 19] indicating a twist of the C-5 methine bridge due to steric hindrance of the asymmetrically substituted ring A with rings B and D. Accordingly, the P-helical form would be favoured, in agreement with both the sign and the strong optical activity of denatured PC.

This interpretation likewise accounts for both the low optical activity observed for denatured PE (chromophore 4) [6] and the large $\Delta \varepsilon$ -values reported for denatured red-absorbing phytochrome (Pr, chromophore 2) [7]. The chromophores of P_r (2) and PC (1) are constitutionally and configurationally closely related. The reason for the high optical activity in the denatured state of Pr may then likewise be attributed to the influence of the chirality centers at C-2, C-3 and C-3¹, in generating an excess population of the P-type tetra-pyrrolhelix. In PE (4), the additional chirality center at C-16 leads to a pertubation at the C-15 methine bridge as well. However, this influence counteracts the one excerted by C-2, C-3 and C- 3^{1} . In a situation similar to that in the i(=inactive)-urobilins with asymmetric C-4 and C-16 [20] these opposing effects in PE account for the extremly low optical activity observed for denatured PE [6].

For a more relevant comparison of PC with P_r we have also measured a cd-spectrum of the ureaunfolded PC under acidic conditions (Fig. 3). The cd-spectrum of protonated PC (Fig. 3) possesses the same phenotype as that of protonated P_r . It is positiv in the long wavelength band, negative in the band around 360 nm. Even the $\Delta \varepsilon$ -values are very similar: P_r: 660 nm, $\Delta \varepsilon = +20$; 375 nm, $\Delta \varepsilon = -15$; PC (values for *one* chromophore): 658 nm, $\Delta \varepsilon =$ + 101/3 = + 33; 350 nm, $\Delta \varepsilon = -75/3 = -25$. In the native state, by contrast, the cd-spectra of phytochrome P_r [7] and of PC (Fig. 1) are (expectedly) quite different indicating the varying influence of the two different proteins on the chromophores.

The close resemblance of the cd-spectra of the denatured PC (Fig. 2, 3) and phytochrome in the P_r-form [7] with that of a model pigment, 2,3-dihydrooctaethylbilindion (3), in chiral, optically active solvents [17] further corroborates the arguments made above. The cd-maxima of the latter at 610/348 nm [17] are only slightly shifted as compared to denatured PC at 615/358 nm (Fig. 2). Moreover, the similar ratios of $\Delta \varepsilon$ -values (0.7 for dihydrooctaethylbilindion 3 and 0.8 for PC in 8 M urea) indicate a similar pitch for the helix of the two chromophores.

c. Stepwise unfolding of PC during denaturation

In an earlier study of PC unfolding [5] an intermediate during the denaturation process of PC with urea has been postulated to account for the singular behavior of the UV-Vis absorption spectra. Fig. 5 nicely shows that the cd-titration curves yield more direct evidence for this assumption. The titration curve of the apoprotein ($\lambda = 220$ nm, curve **a** in Fig. 5) taken from the data of Fig. 4 has an inflection point between 4 M and 5 M urea. This point roughly represents the "melting point" of the apoprotein of PC. However, monitoring the chromophoric region by cd, e.g. at 380 nm and 610 nm (curves **b** and **c** in Fig. 5), curves are obtained whose maxima approximately coincide with the inflection point of curve a for the peptide region (arrows). In a mathematical sense, curves **b** and **c** are derivatives of a. This means, that at least three species of PC are involved in the denaturation process: Between 0 M and 2 M urea the native species dominates. With increasing urea concentration, the concentration of this species decreases and an intermediate, new species appears reaching a maximum at the "melting point" of the apoprotein (4 M - 5 M urea). Higher urea concentrations then favour the completely randomized, denatured PC and the concentration of the intermediate vanishes again at 8 M urea.

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H. Lehner and H. Scheer · Circulardichroism of C-Phycocyanin

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