Receptor and $\beta\gamma$ Binding Sites in the α Subunit of the Retinal G Protein Transducin

René Onrust,* Paul Herzmark, Patty Chi, Pablo D. Garcia, Olivier Lichtarge, Chris Kingsley, Henry R. Bournet

Transmembrane receptors for hormones, neurotransmitters, light, and odorants mediate their cellular effects by activating heterotrimeric guanine nucleotide-binding proteins (G proteins). Crystal structures have revealed contact surfaces between G protein subunits, but not the surfaces or molecular mechanism through which $G\alpha\beta\gamma$ responds to activation by transmembrane receptors. Such a surface was identified from the results of testing 100 mutant α subunits of the retinal G protein transducin for their ability to interact with rhodopsin. Sites at which alanine substitutions impaired this interaction mapped to two distinct G_{α} surfaces: a β_{γ} -binding surface and a putative receptor-interacting surface. On the basis of these results a mechanism for receptor-catalyzed exchange of guanosine diphosphate for guanosine triphosphate is proposed.

Heterotrimeric G proteins relay signals from seven-transmembrane-spanning receptors (7TMRs) to cellular enzymes and ion channels. Activated by photons, odors, and many hormones and neurotransmitters, 7TMRs catalyze replacement by guanosine triphosphate (GTP) of guanosine diphosphate (GDP) bound to $G\alpha$ subunits, causing dissociation of the Ga-GTP and $\beta\gamma$ subunits, which in turn relay the signal to downstream effectors (1). Activation is robust; in the case of transducin (G,), rhodopsin induces an $\sim 10^7$ -fold increase in the basal (unstimulated) rate of GTP-GDP exchange (2). Crystal structures (3-5) have defined conserved folds of G proteins, as well as the trimeric structure, GTP-induced conformational change, and a plausible catalytic mechanism for GTP hydrolysis, which turns off the signal. It is not known, however, how the G protein interacts with its 7TMR and responds by exchanging GDP for GTP.

Biochemical and molecular genetic approaches have defined three key features of this interaction: (i) The 7TMR interacts directly with both $\beta\gamma$ and $G\alpha$ (6, 7). (ii) The COOH-terminal tail of $G\alpha$ (~10 residues) interacts directly with the receptor (7-10). (iii) In addition to the COOHterminal tail, the interaction involves other residues in G α not yet identified (7, 9). Here we report results of a comprehensive molecular genetic strategy that confirm these findings and identify a putative 7TMR-interacting surface of $G\alpha$, and we

Department of Cellular and Molecular Pharmacology, Department of Medicine, Programs in Cell Biology and Biomedical Sciences, and the Cardiovascular Research Institute, University of California, San Francisco, CA 94143-0450, USA

*Present address: Genesis Research and Development, Post Office Box 50, Auckland, New Zealand, †To whom correspondence should be addressed at

S-1212, Box 0450, University of California Medical Center, San Francisco, CA 94143-0450, USA.

suggest a mechanism for 7TMR-triggered conformational change.

We used two assays of G protein activa-

tion to test 100 Ga mutants in which alanine replaced individual amino acid residues in the α subunit of $G_t(\alpha_t)$ [92 single and 8 double replacements (11)]. In the crystal structure, almost all of these residues are oriented toward the solvent (12). The first assay measured the ability of rhodopsin, in the presence of all-trans-retinal, to promote guanosine 5'-O-(3'-triotriphosphate) (GTP- γ -S) binding to α_t in microsomal extracts from COS-7 cells transiently expressing α_t , β_1 , γ_1 , and opsin. Binding of GTP- γ -S induces a conformational change that is detected by the accumulation of a trypsin-resistant α_r fragment (Fig. 1) (10). The second assay (Fig. 2) assessed the ability of α_t , translated in vitro and labeled with [35S]methionine, to bind photoactivated rhodopsin in membranes prepared from retinal rod outer segments (13). Activation by AlF₄⁻ served as a functional control in both assays.

The microsomal assay revealed unequiv-



sentative mutant a, molecules by rho-Microsomal fractions from COS-7 cells expressing α_{t} , alanine mutants, bovine

opsin, β₁, and γ1 were incubated with all-trans-retinal under ambient light. Samples were removed at the indicated times before and after the addition of GTP-y-S (10 µM) and treated with trypsin, as described (10). Lanes labeled "-Retinal" show the slow rate of GTP-y-S-for-GDP exchange in the absence of photoactivated rhodopsin. Lanes labeled "AIF₄⁻⁻" represent incubations that contained AIF₄⁻⁻, but not GTP- γ -S (10). Arrows indicate the 31-kD trypsin-resistant α_t fragments. Standard lanes represent percentages of the original microsomal extract, not treated with trypsin. Abbreviations: WT, normal a; mRR, moderately impaired receptor response; sRR, severely impaired receptor response; WT*, normal interaction when assay results are corrected for a moderate decrease in protection by AIF, I, indeterminable activation by rhodopsin. For the mutant designations, A is Ala, I is Ile, K is Lys, and Y is Tyr, and the number indicates the position of the mutated residue. (B) Quantitative analysis (by densitometry of immunoblots) of α , protection by activated rhodopsin (R*) or AIF₄⁻. For each mutant, values represent the mean ± 2 SEM of at least four separate experiments, performed on microsomal membranes derived from at least two separate transfections. The shaded area depicts the range (mean ± 2 SEM) of results with normal a, (WT) in the presence of all-trans-retinal. (C) Values of activation time courses, corrected for the amount of protection by AIF_4^{-1} [calculated as the quotient of the uncorrected values divided by the mean values for AIF₄⁻ protection (11)]

SCIENCE • VOL. 275 • 17 JANUARY 1997

ocally impaired responses to rhodopsin in 24 of the 88 mutants (11) in which the effect of rhodopsin could be determined. We subclassified these (11) as severely or moderately impaired in receptor response (sRR or mRR); Fig. 1 depicts a representative sRR and an mRR phenotype. The remaining 64 mutants showed normal (WT) responses. Of the 54 in vitro-translated α_r mutants in which we could assess binding to photoactivated rhodopsin, 22 had impaired receptor binding (RB), whereas 32 phenotypes were WT. Autoradiograms and quantitation (by PhosphorImager) of α_{t} bound to rhodopsin, in the absence or presence of GTP- γ -S, for four representative mutants is shown in Fig. 2. In the presence of GTP- γ -S, considerably smaller amounts of [³⁵S]methionine-labeled normal (unmutated) α_{r} bound to rhodopsin, presumably because rhodopsin efficiently induces α_r to bind GTP- γ -S, putting it into a stable conformation that binds poorly to both $\beta\gamma$ and to rhodopsin. For all mutants, classification into WT or RB categories was the same whether we assessed total α_r binding or the difference between binding in the absence or presence of GTP- γ -S (14).

As a control in both assays, α_t mutants were tested for susceptibility to protection

from tryptic cleavage by a receptor-independent stimulus, AlF_4^- [which mimics the γ -phosphate of GTP, thereby switching GDP-bound G α into a conformation that resists trypsin cleavage (3, 4, 10, 15)]. For some mutants in both assays, AlF_4^- protection was reduced but reproducible. In these cases we assessed the rhodopsin interaction phenotype by normalizing it relative to the degree of protection effected by AlF_4^- (Figs. 1 and 2, mutants I339A and D311A, respectively). With the normalization procedure we classified 15 and 7 mutant phenotypes as WT in the activation and binding assays, respectively. By the criterion of resistance to trypsin, small numbers of mutants failed to respond to AlF_4^- under each assay condition (12 of 100 and 12 of 66 in the microsomal and in vitro translation assays, respectively) (16). Rhodopsin activation or binding could not be assessed for these mutants, which are designated (11) indeterminable (I) (Y316A in Fig. 1, L349A in Fig. 2). The phenotypes of all but seven mutants could be determined in at least one of the assays (16). The informative 93 mutants were thus designated either R or WT, to indicate whether or not the mutation impaired interaction with rhodopsin (11).



Fig. 2. Binding to photoactivated rhodopsin of normal and mutant α_t translated in vitro. (**A**) Samples of the diluted translation mix, containing [³⁵S]methionine-labeled α_t , were incubated with rod outer segment membranes in ambient light, in the presence or absence of GTP- γ -S, as described (13). Standard lanes represent percentages of total labeled α_t in the translation extract. Samples were subjected to SDS–polyacrylamide gel electrophoresis (SDS-PAGE) and autoradiography (13). For the mutant designations, D is Asp and L is Leu. (**B**) Quantitative analysis (by PhosphorImager) of binding results. R* binding depicts the percentage of total radiolabeled α_t bound to rhodopsin in the absence (-) or presence (+) of 100 μ M GTP- γ -S. For each mutant, values represent the mean ± 2 SEM of at least four separate experiments, performed on in vitro–translated α_t derived from at least two separate preparations. (**C**) Autoradiograms and quantitative analysis of the percent of α_t protected from trypsin by AlF₄⁻; arrows indicate the 31-kD trypsin-resistant α_t fragments. Std., standard (100% equivalent). (**D**) Corrected values of binding experiments for representative alanine mutants. Bars represent the difference between binding in the absence and the presence of GTP- γ -S, normalized with respect to protection by AlF₄⁻ (11).

To construct a three-dimensional (3D) model of G protein activation by 7TMRs (Fig. 3), we assigned to 34 R residues a specific role as components of binding surfaces that may interact directly with $\beta\gamma$ or rhodopsin. Although any alanine substitution could indirectly hinder the interaction by blocking rhodopsin-induced conformational change, rather than by preventing binding of α_t to rhodopsin, assignments of several residues into the R category are in keeping with previous evidence (5, 7, 9, 17).

Mapped onto the crystal structure (5) of α_t , these 34 R residues form two sets of clusters. Our model assigns these to putative surfaces for interaction with $\beta\gamma$ or with the 7TMR (Fig. 3, A through D). Both surfaces of α_t are probably similar in location and function to corresponding surfaces of other G α molecules (1, 4, 5, 7). Indeed, the two surfaces functionally defined in our experiments closely match surfaces predicted (18) by inference from mapping residues conserved among G α subunits onto the α_t 3D structure.

Rhodopsin and other 7TMRs appear to "act at a distance" (18) in catalyzing GTPfor-GDP exchange, in that intracellular loops of most 7TMRs are too short to allow direct interaction with the guanine nucleotide-binding site, which is probably located ~30 Å from the plasma membrane (Fig. 3, A and C). On the basis of the proposed $\beta\gamma$ - and 7TMR-interacting surfaces, we devised an explanation for propagation of the conformational signal from the receptor to the nucleotide-binding pocket.

Most of the α_t residues that directly contact $\beta\gamma$ in crystals of the heterotrimer (5) showed an R phenotype (17), confirming that the 17 R residues (cyan-colored in Fig. 3, A through C) coincide with the $\beta\gamma$ -binding surface of α_t . $\beta\gamma$ -interacting residues are distributed among four elements of secondary structure (11), three of which (Fig. 3, D and E) stretch from the postulated plane of the plasma membrane (5, 18) to the GDP-binding pocket. It is likely that $\beta\gamma$ cooperates with the 7TMR to open the nucleotide-binding site, either actively, by inducing conformational change in $G\alpha$, or indirectly, by enhancing affinity of the $G\alpha\beta\gamma$ complex for the 7TMR; in the latter role, $\beta\gamma$ could also serve as a stabilizer or fulcrum to make $G\alpha$ more susceptible to a separate action of the 7TMR.

A putative 7TMR-interacting surface (outlined in dark blue in Fig. 3) is located near the membrane, at the ends of $\alpha 5$ and $\beta 6$ (that is, the COOH-terminal tail and the $\alpha 4$ - $\beta 6$ loop, respectively; Fig. 3, A through C). Additional mutation-sensitive sites located in both $\beta 6$ and $\alpha 5$ extend part way toward the guanine nucleotide-bindFig. 3. Alanine mutations and 3D structure of α_t . (A to C) Views of at rotated successively by 90° about the vertical axis, which is parallel to the postulated (5, 18) plane of the plasma membrane; the membrane presumably is located just to the right of the model in (A) and just to the left of the model in (C) (vertical yellow stripes). Panel (A) represents the face of α_{t} that binds to $\beta\gamma$ and (B) the face of α_{t} that would be "seen" by the receptor in the membrane, whereas (C) depicts the face of α_{+} opposite to that seen in (A). Residues at sites where alanine substitution impaired interaction with rhodopsin are colored cyan (βγ-interacting surface, outlined in magenta), red (putative rhodopsin-interacting surface, outlined in dark blue), and green [other residues, as indicated in (24)]. Beige or black indicate, respectively, sites at which mutations produced WT or indeterminate phenotypes. In (A) and (C), GDP (yellow) can be glimpsed between the two $\mbox{G}\alpha$ domains (the α -helical domain is colored gray). Ct, COOH-terminal residues 344 to 350 of α_{t} ; not visible in crystals of the heterotrimer (5), these are taken from a separate crystal structure (3). Nt, NH2-terminus. (D and E) Ribbon representations of α_t in a complex (5) with subunit β (dark blue; γ not



shown). In (D) and (E) α_t is tilted ${\sim}90^\circ$ around the horizontal axis with respect to (A), so that the $\dot{NH}_2\text{-terminal }\alpha\text{-helix}$ points toward the viewer. Panel (D) shows locations of sites with an R phenotype, colored as in (A) to (C). Panel (E) highlights structural elements postulated to transmit conformational change from the receptor to the guanine nucleotide-binding

pocket. These include $\alpha 2$ and $\beta 1$ through $\beta 3$ (cyan), which communicate with $\beta\gamma$, and $\beta6$ and $\alpha5$ (red), proposed as key conformation transmitters between the $\beta 6-\alpha 5$ loop (green, contacting GDP) and parts of the protein that interact directly with membrane-bound receptor, R (a4-B6 loop, midα5, and the COOH-terminus).

ing pocket (11). We propose (Fig. 3, D and E) that 7TMR-induced changes in the conformations of the $\beta 6$ strand and the $\alpha 5$ helix are communicated to the β 6- α 5 loop (green in Fig. 3E), in which side chains of a short sequence of amino acids contact the guanine ring of GDP (3, 4, 18, 19).

Although not yet explicitly tested, this idea is in keeping with our present observations and with results of studies that probed functions of $\beta 6$, $\alpha 5$, and the $\beta 6$ - $\alpha 5$ loop (20-23). Specifically, several mutations in $\alpha 5$, at locations distant from the nucleotide-binding pocket, increase rates of GDP dissociation and its replacement by GTP (20). Amino acid substitutions in the $\beta 6-\alpha 5$ loop increase the GDP dissociation rate in several guanosine triphosphatases (GTPases), promoting spontaneous GTP-for-GDP exchange (21-23). Such mutations dramatically activate both p21^{ras} and α_s —producing, respectively, neoplastic transformation (22) and a human endocrine syndrome, testotoxicosispseudohypoparathyroidism (23). Thus 7TMR-induced conformational change may be propagated from the membrane through $\beta 6$ and $\alpha 5$ to the loop that connects them, right in the GDP-binding pocket (Fig. 3, D and E).

REFERENCES AND NOTES

- 1. H. R. Bourne, D. A. Sanders, F. McCormick, Nature 348. 125 (1990); L. Birnbaumer, Annu. Rev. Pharmacol. Toxicol. 30, 675 (1990); A. H. Cheuna, R. R. Huang, C. D. Strader, Mol. Pharmacol. 41, 1061 (1992); M. Freissmuth, P. J. Casey, A. G. Gilman, FASEB J. 3, 2125 (1989); Y. Kaziro, H. Itoh, T. Kozasa, M. Nakafuku, T. Satoh, Annu. Rev. Biochem. 60, 349 (1991); E. J. Neer, Cell 80, 249 (1995).
- T. M. Vuong, M. Chabre, L. Stryer, *Nature* **311**, 659 (1984); L. Ramdas, R. M. Disher, T. G. Wensel, *Bio*chemistry 30, 11637 (1991).
- 3. J. P. Noel, H. E. Hamm, P. B. Sigler, Nature 366, 654 (1993).
- 4. D. G. Lambright, J. P. Noel, H. E. Hamm, P. B. Sigler, ibid. 369, 621 (1994); J. Sondek, D. G. Lambright, J. P. Noel, H. E. Hamm, P. B. Sigler, ibid. 372, 276 (1994); D. E. Coleman et al., Science 265, 1405

(1994); M. A. Wall et al., Cell 83, 1047 (1995).

- D. G. Lambright *et al.*, *Nature* **379**, 311 (1996).
 B. K.-K. Fung and C. R. Nash, *J. Biol. Chem.* **258**, 10503 (1983); W. J. Phillips and R. A. Cerione, ibid. 267, 17032 (1992); W. J. Phillips, S. C. Wong, R. A. Cerione, *ibid.*, p. 17040; D. J. Kelleher and G. L. Johnson, *Mol. Pharmacol.* **34**, 452 (1988); O. Kisselev and N. Gautam, J. Biol. Chem. 268, 24519 (1993); A. B. Fawzi et al., ibid. 266, 12194 (1991).
- 7. Reviewed in B. R. Conklin and H. R. Bourne, Cell 73, 631 (1993)
- 8. K. A. Sullivan et al., Nature 330, 758 (1987); E. A. Dratz et al., ibid. 363, 276 (1993).
- H. E. Hamm et al., Science 241, 832 (1988).
- P. D. Garcia, R. Onrust, S. M. Bell, T. P. Sakmar, H. R. Bourne, EMBO J. 14, 4460 (1995)
- A supplementary table, available with the Science Online version of this report at http://www. sciencemag.org/ lists each of the individual residues mutated, along with their phenotypes, in both rhodopsin interaction assays. The supplement also details the citeria we used to assign WT and R "diagnoses" to each mutant and the procedure for using the AIF₄ - data for normalizing results of the rhodop sin interaction assays. We have reported results of the microsomal assay, but not the in vitro translation assay, for the 10 mutants in the COOH-terminal tail of a, (10).
- 12. Sites were chosen for mutation if previous work suggested involvement of a particular structural element

in receptor interaction [for instance, the COOH-terminus, $\alpha 4$ - $\beta 6$ loop, and $\beta 6$ (7-10)], if they were known to interact with $\beta\gamma$ in the 3D structure of $\alpha_+(5)$, or if they were located on the surface of the 3D structure of a + near either known or postulated rhodopsin- or βy-interacting residues (3-5)

- 13. S. Osawa and E. R. Weiss, J. Biol. Chem. 270, 31052 (1995).
- 14. That is, binding values for different mutants in the presence of GTP- γ -S were consistent and small enough not to confound the results. We failed to find a mutant α_t that could bind to rhodopsin but would not be susceptible to rhodopsin-catalyzed replacement of GDP by GTP-y-S. A mutant rhodopsin with this phenotype has been described (O. P. Ernst, K. P. Hofmann, T. P. Sakmar, ibid., p. 10580).
- J. Bigay, P. Deterre, C. Pfister, M. Chabre, *FEBS Lett.* **191**, 181 (1985); P. C. Sternweis and A. G. 15. Gilman, Proc. Natl. Acad. Sci. U.S.A. 79, 4888 (1982)
- 16. The failure of individual mutants to be protected from trypsin by AIF4- could indicate improper folding of the protein or an inability of the α2 helix, in which the protected cleavage site is located, to take on the trypsin-resistant (active, GTP-bound) conformation. Of the 66 mutants tested in both assays, only four showed an I phenotype in both. It is likely that overexpression of recombinant α_t in COS cells and its

translation in vitro affect folding in different ways.

- 17 Of the 23 residues identified (5) as interacting directly with βγ, we tested 22. By one or the other assay, mutational replacement of 14 of the 22 residues (in 10 mutants) produced an R phenotype, whereas six mutants were WT and two were indeterminable (11).
- 18. O. Lichtarge, H. R. Bourne, F. E. Cohen, Proc. Natl. Acad. Sci. U.S.A. 93, 7507 (1996). In this theoretical study, residues in cluster 2 and cluster 1 correspond, respectively, to the \$\beta\gamma- and putative 7TMR-interacting surfaces identified in the present work.
- 19. This loop comprises residues 321 to 324 of α_{t} . Corresponding \$6-a5 loops of other GTPases interact similarly with the guanine ring of bound GDP and GTP [H. R. Bourne, D. A. Sanders, F. McCormick, Nature 349, 117 (1991)].
- B. M. Denker, C. J. Schmidt, E. J. Neer, J. Biol. 20 Chem. 267, 9998 (1992); B. M. Denker, P. M. Bou-
- tin, E. J. Neer, Biochemistry 34, 5544 (1995); L. A. Quilliam et al., Proc. Natl. Acad. Sci. U.S.A. 92, 1272 (1995).
- T. C. Thomas, C. J. Schmidt, E. J. Neer, Proc. Natl. 21 Acad. Sci. U.S.A. 90, 10295 (1993).
- 22 L. A. Feig and G. M. Cooper, Mol. Cell. Biol. 8, 2472 (1988). T. liri, P. Herzmark, J. M. Nakamoto, C. Van Dop, H.
- 23 R. Bourne, Nature 371, 164 (1994).
- 24 An indirect effect of alanine substitution is especially likely for the few residues we tested whose side

Consumer Versus Resource Control in Freshwater Pelagic Food Webs

Michael T. Brett* and Charles R. Goldman

Models predict that food-web structure is regulated by both consumers and resources, and the strength of this control is dependent on trophic position and food-web length. To test these hypotheses, a meta-analysis was conducted of 11 fish (consumer)-bynutrient (resource) factorial plankton community experiments. As predicted, zooplankton biomass was under strong consumer control but was weakly stimulated by nutrient additions; phytoplankton biomass was under strong resource control with moderate control by fish. However, the phytoplankton and zooplankton responses to nutrient additions did not follow theoretical predictions based on the number of trophic levels in the food web.

 ${f T}$ he nature of the factors regulating foodweb structure has been a very active area of ecological research (1, 2) since the classic paper by Hairston, Smith, and Slobodkin (3) was published in 1960. In aquatic systems, food-web interactions strongly influence fisheries production, biogeochemical cycling, and ecosystem responses to anthropogenic eutrophication. A recent quantitative summary of the freshwater trophic cascade (4) literature showed that planktivorous fish treatments result in decreased herbivore (zooplankton) and increased primary producer (phytoplankton) biomass (5). In addition, phytoplankton response to the cascade is weakly dampened and highly variable, with weak responses in two-thirds of the experiments and very strong responses in the other experiments (5). Still, many questions regarding the dynamic nature of food-web interactions remain unresolved (1, 2). In particular, what is the relative strength of consumer and resource control in pelagic food webs (6), and how do food webs respond to changes in system productivity under different food-web configurations (7)?

The debate over top-down (consumer) versus bottom-up (resource) control represents a synthesis of the known impact of nutrient regulation of primary producers (8) and higher trophic levels (9), and the more recent emphasis on consumer control of trophic levels through the cascade (4). In essence, the debate centers on whether herbivore and plant communities are regulated through predator control of herbivore abundance or through nutrient control of primary production. McQueen and colleagues (6) predicted bottom-up control is stronger at the base of the food web, and top-down control is stronger at higher trophic levels.

chains are directed toward the protein's hydrophobic core; these include two phenylalanine residues, colored cyan (F185) or red (F332) in Fig. 3, A through C. We considered four additional R residues (green in Fig. 3, A through C), scattered through the α_{+} molecule, as quite unlikely to interact directly with $\beta\gamma$ or rhodopsin. Residues Gly² and Ser⁶ are required for myristoylation of $\alpha_{\rm p}$, which in turn is necessary for efficient activation [P. J. Casey, *Curr. Opin. Cell Biol.* **6**, 219 (1994)]. Residues Thr³²³ and Asp³²⁴ are located very close to the guanine nucleotide-binding pocket (3-5), where they might be expected to alter regulation of GTP-GDP exchange.

25. We thank members of our laboratories for helpful discussions and reading of the manuscript, H. Czerwonka for preparing the manuscript, H. Hamm (University of Illinois at Chicago) for a sample of rod outer segment membranes, and B. K.-K. Fung (Jules Stein Eye Research Institute, Los Angeles, CA) for the TF15 monoclonal antibody used for α_{t} immunoblots. R.O. is a Helen Hay Whitney fellow, P.C. was supported by a grant from the Genentech Foundation, P.D.G. was supported in part by funds from Daiichi, and O.L. was supported by funds from the American Heart Association. Supported by NIH grants GM-27800 and CA-54427 (to H.R.B.).

27 August 1996; accepted 12 November 1996

For example, the zooplankton should be more strongly controlled by zooplanktivorous fish than by nutrients, whereas phytoplankton biomass should be primarily controlled by nutrient availability and to a lesser extent by higher trophic levels.

Oksanen et al. (7) developed a series of models to explore the theoretical relationship among ecosystem productivity, patterns of biomass accrual, and the number of trophic levels in that ecosystem. This predicted "a stepped pattern of biomass accrual" (2) across productivity gradients (10). In food webs with an odd number of trophic levels, increases in primary production should lead to increased biomass for oddnumbered trophic levels and no change in biomass for even-numbered trophic levels. Conversely, in food webs with an even number of trophic levels, increases in primary production should lead to increased biomass for even-numbered trophic levels and no change in biomass for odd-numbered trophic levels.

We assembled eight studies (11) that reported the results of 11 independent mesocosm experiments employing factorial nutrient addition and zooplanktivorous fish treatments. Simple criteria were used to decide which studies to include in our analvsis (12). Six of the studies used simple fish-by-nutrient designs, and two used slight modifications of this design (13). In five studies, zooplankton community biomass values were obtained directly, and in three studies, zooplankton biomass was estimated using abundance and individual biomass data (14). All phytoplankton community biomass values were taken directly from the respective studies.

Mesocosms are classic experimental de-

Division of Environmental Studies, University of California, Davis, CA 95616, USA.

^{*}To whom correspondence should be addressed. E-mail: mtbrett@ucdavis.edu