Lateral Hypothalamic Signaling Mechanisms Underlying Feeding Stimulation: Differential Contributions of Src Family Tyrosine Kinases to Feeding Triggered Either by NMDA Injection or by Food Deprivation

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In rats, feeding can be triggered experimentally using many approaches. Included among these are (1) food deprivation and (2) acute microinjection of the neurotransmitter L-glutamate (Glu) or its receptor agonist NMDA into the lateral hypothalamic area (LHA). Under both paradigms, the NMDA receptor (NMDA-R) within the LHA appears critically involved in transferring signals encoded by Glu to stimulate feeding. However, the intracellular mechanisms underlying this signal transfer are unknown. Because protein-tyrosine kinases (PTKs) participate in NMDA-R signaling mechanisms, we determined PTK involvement in LHA mechanisms underlying both types of feeding stimulation through food intake and biochemical measurements. LHA injections of PTK inhibitors significantly suppressed feeding elicited by LHA NMDA injection (up to 69%) but only mildly suppressed deprivation feeding (24%), suggesting that PTKs may be less critical for signals underlying this feeding behavior. Conversely, food deprivation but not NMDA injection produced marked increases in apparent activity for Src PTKs and in the expression of Pyk2, an Src-activating PTK. When considered together, the behavioral and biochemical results demonstrate that, although it is easier to suppress NMDA-elicited feeding by PTK inhibitors, food deprivation readily drives PTK activity in vivo. The latter result may reflect greater PTK recruitment by neurotransmitter receptors, distinct from the NMDA-R, that are activated during deprivation-elicited but not NMDA-elicited feeding. These results also demonstrate how the use of only one feeding stimulation paradigm may fail to reveal the true contributions of signaling molecules to pathways underlying feeding behavior in vivo.

**Key words:** eating; feeding; lateral hypothalamic area; glutamate; NMDA; NR2A; NR2B; Src; Pyk2; tyrosine phosphorylation; signal transduction

**Introduction**

In recent years, molecular and cellular techniques have helped identify novel macromolecules involved in feeding control and energy homeostasis. Ironically, however, the molecular and cellular mechanisms underlying the actions of these agents remain poorly understood in the freely behaving animal. Identifying such mechanisms is important, especially for neurons within various hypothalamic nuclei, where neuropeptides and small neurotransmitters implicated in feeding control may act on common neural substrates. In such neurons, a few intracellular effectors could integrate multiple signals encoding information important for feeding control. Identifying such effectors could reveal novel targets for treating certain eating and metabolic disorders or cases of drug dependency (Carr, 1996, 2002; Chiesi et al., 2001; DiLeone et al., 2003). Despite progress in identifying mammalian feeding control circuits, the intracellular pathways they use remain virtually unknown in vivo, especially for feeding stimulation.

This is partly attributable to limitations in methods. In the rat, which has the best delineated feeding control neural circuitry but for which a recently developed gene knock-out strategy (Zan et al., 2003) has yet to gain widespread currency (Rossant, 2003), approaches in vivo have primarily involved (1) administering central and peripheral agents that target signaling molecules to alter feeding behavior and (2) manipulating the behavioral state (e.g., food deprivation) to change signaling molecule activity. The former approach poorly reflects the concerted physiological activity of neurochemical systems in vivo, and the latter rarely identifies unambiguously the causal role of a particular signaling

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**Behavioral/Systems/Cognitive**

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molecule or pathway. However, combining these approaches could mitigate the limitations attendant with each, revealing more accurately the intracellular mechanisms engaged by various feeding-related neurochemical systems.

One such system is that for the amino acid l-glutamate (Glu) in the lateral hypothalamic area (LHA) (Stanley, 1996). In satiated rats, LHA-administered Glu or its receptor agonists (kainate and NMDA), elicit powerful, transient feeding behaviors that are stereochemically, anatomically, and behaviorally specific (Stanley et al., 1993b,c; Duva et al., 2001, 2002). Conversely, LHA-injected NMDA receptor (NMDA-R) antagonists suppress eating elicited by NMDA, food deprivation, or the onset of the nocturnal period and reduce body weight when given chronically (Stanley et al., 1996, 1997; Khan et al., 1999). During fasting, endogenous Glu is apparently released within the LHA during meal onset, sharply declining while rats are still eating (Rada et al., 1997, 2003). The orexin hypocretin/orexin (H/O) requires Glu signals to depolarize H/O-containing neurons within the LHA (Li et al., 2002), and neuropetide Y requires functional LHA NMDA-Rs to elicit feeding (Lee and Stanley, 2002). Thus, LHA signaling mechanisms underlying feeding behavior are probably driven by multiple neurotransmitters and their receptors, including Glu and its NMDA-R subtype.

Accordingly, we wanted to identify LHA signaling pathways activated during the feeding stimulated either by NMDA injection, or more generally, by food deprivation. Here we present evidence suggesting that feeding stimulation is associated with the robust activation of LHA protein-tyrosine kinases (PTKs), which are also required for the full expression of feeding behavior in vivo.

Parts of this paper have been published previously in preliminary form (Khan et al., 1996, 2000a, 2001; Khan, 2002).

### Materials and Methods

#### Central injection studies

**Reagents and solutions for central injection studies**

Genistein [4’7,5,7-trihydroxysflavone; molecular weight (MW) 270.2], daidzein [4’7-dihydroxysflavone; MW 254.2], and tyrphostin A48 [also known as AG112, 2-amino-4-[(4’-hydroxyphenyl)-1,1,3-tricyanobuta-1,3-diene; MW 236.2] were purchased from Calbiochem (La Jolla, CA). PP1 [4-amino-5-(4-methylphenyl)-7-(2-buty1)pyrazolo[3,4-d]pyrimidine; MW 281.4] was obtained from Biomol (Plymouth Meeting, PA). The PTK inhibitors were dissolved in dimethylsulfoxide (DMSO; Sigma, St. Louis, MO), a vehicle recently validated as useful in studies of chemically elicted feeding in the hypothalamus (Blevins et al., 2002). The light-sensitive genistein and daidzein solutions were prepared in dim light, and vials of working solutions were covered with aluminum foil. NMDA and kainic acid (KA) were also purchased from Sigma and were dissolved in artificial CSF (aCSF) just before central injection. Sterile aCSF (in msc 147 Na+, 154.2 Cl−, 3.0 K+, 1.2 Ca2+, and 0.9 Mg2+, pH 7.4) was prepared using double-distilled water and subsequently filtered through 0.22 μm GS-type or 0.45 μm HV-type filters (Millipore, Bedford, MA). For the experiments using PP1, the aCSF was also buffered with 1.0 mM potassium phosphate. Materials for histological verification of cannula placement were purchased from Sigma.

**Central injection methods**

**Subjects.** Adult male Sprague Dawley rats (n = 127; 350–500 gm), descended from Charles River Laboratories (Wilmington, MA) animals, were single-housed in a temperature-controlled vivarium with a 12 h light/dark photoperiod and ad libitum access to food and water, unless otherwise stated. During testing, animals were maintained on a mash diet consisting of 46% laboratory rodent meal (LabDiet; PMI Nutrition International), 37% sucrose, and 17% evaporated milk. Although we used this mash diet instead of chow pellets because it minimizes spillage and therefore helps us measure intake more accurately, rats given LHA NMDA injections will also readily eat normal chow pellets (A. Khan, unpublished observations). Except for food deprivation studies, animals were fed freshly prepared mash at least 1.5 hr before testing to maximize satiety.

**Stereotaxic surgery.** Animals in central injection studies were stereotaxically implanted under methoxyflurane inhalant (Metofane; Schering-Plough, Kenilworth, NJ) or pentobarbital anesthesia (50 mg/kg, i.p.) with chronic unilateral or bilateral 26 gauge stainless steel indwelling guide cannulas (Small Parts, Miami Lakes, FL) targeted toward the LHA. With the incisor bar 3.3 mm below the ear bars, stereotaxic coordinates were 6.1–6.4 mm anterior to the interaural line, 1.8 mm lateral to the mid sagittal sinus, and 8.2 mm ventral to the surface of the skull (Paxinos and Watson, 1998, their Figs. 30, 31). Cannulas were secured to the skull by six stainless steel hex screws and dental acrylic, and a plastic guard was placed around the exposed portion of the cannula. A 33 gauge stainless steel obturator was inserted into each cannula to maintain patency. Animals were allowed 5–7 d of postoperative recovery before testing, during which time they were repeatedly handled and mock-injected to adapt them to the testing procedures.

**Behavioral testing.** Tests were conducted in the midportion of the light phase of the rat circadian cycle and consisted of either acute bilateral injections (one injection per cannula in quick succession) or acute unilateral injections in series 10 min apart (vehicle, tyrphostin A48 or PP1, followed by vehicle or NMDA) or 30 min apart (vehicle, genistein or daidzein, followed by vehicle, NMDA, or KA). Solutions (300 nl volume) were delivered through 33 gauge injectors, each inserted through the guide cannula and projecting exactly 1.0 mm beyond the end of the cannula, directly into the LHA. Doses of NMDA (10 nmol) and KA (1 nmol) were selected on the basis of their efficacy in eliciting eating in previous studies (Stanley et al., 1993b,c; Khan et al., 1999). Food intake was recorded at 0.5, 1, 2, and 4 hr after the final injection by weighing each subject’s food bowl to the nearest 0.1 gm and subtracting the resulting weight from the weight of the bowl recorded before injection. Animals received treatments in counterbalanced order (Latin square design) during a series of test days, with each animal receiving each treatment by the end of the experiment. Treatment days were separated by at least 1 d of recovery, during which no procedures were performed.

**Testing the effects of genistein on food intake**

**Experiments 1a and 1b: does genistein suppress eating elicited by NMDA?** To determine whether NMDA-elicited eating can be attenuated by the broad-spectrum PTK inhibitor genistein, 24 animals were unilaterally injected in experiment 1a with genistein (240 nmol), its “inactive” analog daidzein (240 nmol), or DMSO vehicle, followed 30 min later by a unilateral injection of NMDA (10 nmol) or aCSF vehicle. In experiment 1b, 18 naive animals were tested in a similar manner, except that three doses of genistein were used (2.4, 24, and 240 nmol), and daidzein was not included as a treatment.

**Experiment 2: does genistein suppress eating elicited by kainic acid?** As with NMDA injection, a unilateral KA injection into the LHA of satiated rats elicits a robust feeding response (Stanley et al., 1993b,c). To test whether genistein could also suppress eating elicited by KA, 15 naive rats received unilateral LHA injections of 240 nmol of genistein or DMSO vehicle, followed 30 min later by LHA injections of either KA (1 nmol) or aCSF vehicle.

**Experiment 3: does genistein suppress eating elicited by food deprivation?** To investigate whether genistein can affect natural eating elicited by food deprivation, 12–15 naive animals with ad libitum access to water were food-deprived for ~20 hr and then given bilateral LHA injections of either genistein (240 nmol/side) or DMSO vehicle. Genistein was given bilaterally in this experiment so that it might block the presumed bilateral activation of the LHA caused by food deprivation. Freshly prepared food was presented 10 min after the injections, and food intake was measured for 60 min of final injection as t = 0.

**Testing the effects of tyrphostin A48 on food intake**

**Experiment 4: does tyrphostin A48 suppress eating elicited by NMDA?** To determine the effect of the PTK inhibitor tyrphostin A48 on NMDA-
elicited eating, 13 naive animals received unilateral LHA injections of one of three doses of tyrphostin A48 (2.1, 21.1, or 211.7 nmol) or DMSO vehicle, followed 10 min later by a unilateral LHA injection of NMDA (10 nmol) or aCSF vehicle.

**Testing the effects of PP1 on food intake**

**Experiments 5a and 5b:** does PP1 suppress eating elicited by NMDA? To help determine the involvement of members of the Src family of PTKs in mechanisms underlying NMDA-elicited eating, 15 naive animals were unilaterally injected in experiment 5a with one of three doses of the Src family-selective inhibitor PP1 (0.2, 2.1, or 21.2 nmol) or DMSO vehicle, followed 10 min later by a unilateral LHA injection of NMDA (10 nmol) or aCSF vehicle. In experiment 5b, this study was replicated using 14 naive rats, except that the 0.2 nmol dose of PP1 was not included.

**Experiment 6:** does PP1 suppress eating elicited by food deprivation? To determine whether Src family PTK activity is important for eating triggered by food deprivation, 13 naive animals were food-deprived under conditions identical to those described in experiment 4 (above), except that the inhibitor, injected bilaterally into the LHA, was PP1 (21.2 nmol/side).

**Statistical and histological analyses for behavioral studies**

After completion of behavioral testing, subjects were killed by CO2 inhalation and then perfused transcardioly with 10% formalin. Using a cryostat, a subset of brains was sectioned into 100-μm-thick sections through the extent of the cannula track. Sections were air-dried on gelatin- or gelatin- and chromate-subbed slides and stained with thionin. Injection sites were localized by tracing their projected images onto size-matched figures adapted from the atlas of Paxinos and Watson (1998). The appearance of the tissue was similar to that of subjects in previous studies (data not shown), the histological photomicrographs of which have been published (Stanley et al., 1993c).

Food intake values were analyzed by one- or two-way general linear model ANOVA. Where overall effects were significant, multiple comparisons were performed using the Student–Newman–Kuels test with α set at 0.05. Nonparametric tests (Mann–Whitney rank sum tests and Kruskal–Wallis ANOVA on ranks) were performed, where appropriate, for scores that were not normally distributed. In such cases, multiple comparisons were performed using Dunn’s test, with α set at 0.05. All food intake scores are expressed as mean intakes (in grams) ± SEM.

**Immunohistochemistry (experiment 7)**

**Reagents and solutions**

The primary polyclonal Src antibody was raised against the gene product of the Rous sarcoma virus, which includes Src kinase (Biomol). Biotinylated goat anti-rabbit IgG (heavy and light chains) and avidin-HRP solution were from Vector Laboratories (Burlingame, CA). All other materials, including bovine serum albumin, hydrogen peroxide, and 3,3’-diaminobenzidine (DAB), tetrachloride tablets were purchased from Sigma, Vector Laboratories, or local suppliers.

**Tissue preparation**

As described previously (Khan et al., 1999, 2000b), adult male Sprague Dawley rats (325–475 gm; n = 11) were anesthetized with sodium pentobarbital (60 mg/kg, i.p.) and perfused through the ascending aorta with at least 100 ml of cold, heparinized PBS (0.01 M sodium phosphate buffer in 0.9% saline, containing 200 U/ml heparin), followed by at least 300 ml of ice-cold 4% paraformaldehyde in 0.1 M sodium phosphate buffer, pH 7.4. The brain was quickly dissected, and a razor blade was used to obtain a block of tissue containing the LHA. These blocks were placed in a 30% sucrose-PBS solution at 4°C for at least 36 hr without an intervening post-fixation step. They were then mounted onto the freezing stage of a sliding microtome using Tissue-Tek or Histomount (Fisher Scientific, Houston, TX), frozen using powdered dry ice, and sliced into 30 μm coronal sections. The sections were then transferred to wells containing cold PBS.

Frothy floating sections were reacted with 0.1% sodium borohydride for 10 min at room temperature (RT), followed by 0.1% hydrogen peroxide in PBS for 10 min at RT, and then rinsed for 10 min in PBS before the blocking step. Blocking involved incubating the sections for 1 hr at RT in 5–10% normal goat serum (NGS) diluted in 0.3% Triton X-100-containing PBS, pH 7.4 (T-PBS).

**Immunodetection**

**Primary and secondary antibody reactions.** Sections were then incubated with the rabbit anti-Src antibody for at least 24 hr at 4°C. Initial experiments were performed using dilutions of 1:500 and 1:1000. Optimal visualization was obtained at a dilution of 1:750. The primary antibody was dissolved in T-PBS containing 0.2% BSA and 1% NGS. Sections were then reacted with a biotinylated goat anti-rabbit IgG (diluted 1:200 in T-PBS) for 1–2 hr at RT.

**Immunoperoxidase detection.** Sections were then treated for 1 hr at RT in avidin-HRP solution (1:200 in T-PBS). They were transferred briefly to a 0.05 M Tris-HCl solution containing 0.9% NaCl, pH 7.4, and incubated in a DAB chromogen solution (20 mg of DAB/100 ml of Tris-HCl) for 5–10 min at RT before being reacted at RT with 1–4 μl of 30% H2O2 in 10 ml of DAB solution. Nickel chloride (0.15 gm/100 ml of DAB solution) was added to the DAB solution to intensify the stain, and reactions were stopped with cold Tris-HCl. The optimized time in the DAB/H2O2 solution was 4 min.

**Immunohistochemical controls**

For a subset of brain sections in all experiments, a solution containing only Triton X-100, serum, and BSA was substituted for the primary antibody.

**Tissue mounting, observation, and photomicrography**

Sections were mounted onto gelatin- and chromate-subbed slides and air-dried. The slides were dehydrated in ascending concentrations of ethanol, cleared in 100% xylene, and coverslipped with DPX. Brain regions were then examined under bright-field illumination using an Olympus Optical (Tokyo, Japan) BH-2 microscope (with an accompanying Olympus EMM-7 light meter) or a Nikon (Melville, NY) Microphot FX-A microscope. In some cases, camera lucida drawings of immunoreactive cells were created by observing sections under oil immersion at 100× magnification. Photomicrographs of immunoreactive regions were prepared using Eastman Kodak (Rochester, NY) Technical Pan film 5612 (ASA 25) or Kodak Gold film (ASA 200). The former was developed in the University of California, Riverside, darkroom facility, where prints were also prepared using Kodak F4 or F5 paper.

**Biochemical identification of tyrosine-phosphorylated LHA proteins**

Experiment 8a refers to biochemical studies performed on tissue from rats receiving injections of NMDA (and their aCSF-injected controls), and experiment 8b refers to similar studies performed on tissue from food-deprived rats (and their ad libitum-fed controls). Unless otherwise noted by experiment number, all procedures described below apply to both experiments.

**Subjects, housing, and treatments**

Adult male Sprague Dawley rats (n = 60), housed under identical conditions as animals used in behavioral experiments, were maintained on a standard pelleted diet (LabDiet) with ad libitum access to water at all times, unless otherwise noted.

**Experiment 8a:** is an acute injection of NMDA into the LHA, which normally triggers a transient eating response, also sufficient to affect Tyr-P of proteins, PTK expression, and PTK activity in LHA-containing tissue? For this experiment, which was conducted between 7 P.M. and 4 A.M. (between the end of the light phase and the beginning of the dark phase), rats (n = 20) each received ~1.0 ml of sodium pentobarbital anesthesia (50 mg/ml, i.p.) to minimize potential trans-synaptic or sensorimotor feed-back effects on tyrosine phosphorylation. When whole-body movements were no longer elicited by a brief tail pinch, rats received central bilateral LHA microinjections of either NMDA (10 nmol; n = 10) or aCSF (n = 10). Parameters for central microinjections (volume and method of delivery) were identical to those described for the behavioral experiments (experiments 1–6). Ten to 15 min after the bilateral LHA injections, each rat was transferred to a cold room for tissue harvesting.

**Experiment 8b:** is prolonged feeding stimulation, as produced by food deprivation, sufficient to affect Tyr-P of proteins, PTK expression, and PTK activity in LHA-containing tissue? For these experiments, which were conducted between 7 P.M. and 12 A.M. (toward the end of the diurnal
phase), four separate tissue harvests were conducted, each on 10 rats, for a total of 40 rats. In each group of 10, 5 rats were food-deprived for 48 hr immediately before the tissue harvest, whereas 5 had ad libitum access to the pellet diet and served as controls. As in experiment 8a, rats received ~1.0 ml of sodium pentobarbital anesthesia (50 mg/ml, i.p.) just before tissue harvest.

Tissue harvest

Each rat was weighed to the nearest 0.1 gm, transferred to a cold room (10°C), and decapitated, and its brain was quickly dissected and immediately immersed in freshly prepared, ice-cold protease inhibitor mixture (PIM), pH 7.4, for 10–15 sec. The PIM contained 0.32 mM sucrose, 50 mM Tris, 0.2 mM sodium orthovanadate (Calbiochem), 20 mM β-glycerophosphate, 1 mM NaF, 0.1 mM phenylmethylsulfonyl fluoride (PMSF), and 5 µg/ml each antipain, leupeptin, and aprotinin (Sigma), all dissolved in double-distilled water. The brain was quickly placed ventral side-up within a chilled stainless steel rat brain mold (Activational Systems) containing grooves 1 mm apart for producing coronal transsections. Chilled PIM was immediately poured over the brain, and coronal transsections were then made. Tissue punches were dissected and incised 10-fold with a razor blade. Diluted the raised surface of the median eminence. These blocks of brain tissue contained virtually all of the LHA and also likely portions of the preoptic and ventral tegmental areas. Each tissue block was placed flat, with the posterior side facing up, onto a chilled microscope slide, and fresh chilled PIM was again poured over it.

With the aid of a dissecting stereoscope (Nikon SMZ-2B) and the atlas of Palkovits and Brownstein (1988), a 14 gauge stainless steel spinal tap catheter was used to core approximately bilaterally symmetrical tissue samples containing the LHA, using the fornix, optic tract, and internal capsule as reference boundaries. Samples were immediately placed in 1.5 ml Eppendorf microcentrifuge tubes containing 1.0 ml of PIM, and tissue wet weight was determined by weighing tared vials to the nearest 0.1 mg. Samples were immediately placed on ice in the cold room and then transferred to a −70°C freezer. They were then shipped overnight, with dry ice, to the University of Toronto at Scarborough, where they were again stored at −70°C until used for immunoprecipitation and immunoblotting.

Immunoprecipitation (experiment 8b only)

As described previously (Takagi et al., 1997; Cheung et al., 2000), NR2A and NR2B antibodies used for immunoprecipitation (IP) were generated against polyhistidine fusion proteins containing the C-terminal region of each subunit corresponding to amino acid residues 934–1203 for NR2A and 935–1455 for NR2B. Antibodies were precoated on Protein A/G PLUS-agarose (Santa Cruz Biotechnology; Santa Cruz, CA) by incubating 0.003 or 0.005 ml of immune serum with 0.02 ml of protein A/G PLUS-agarose for 45 min–1 hr at 4°C in 0.5 ml of 50 mM Tris-HCl buffer, pH 7.4, containing 1% (v/v) NP-40, 0.25% (w/v) sodium deoxycholate, 150 mM NaCl, 1 mM EDTA, 0.1% each PMSF and sodium orthovanadate, 1 mM NaF, and 5 µg/ml each antipain, leupeptin, and aprotinin (IP buffer). Three hundred micrograms of LHA-containing tissue that had been solubilized by boiling in 100 µl of 1% SDS containing 1% β-mercaptoethanol, were diluted 10-fold with binding buffer. Diluted samples were incubated with 0.02 ml of Protein A/G PLUS-agarose for 1 hr at 40°C and centrifuged, and the supernatant was added to the precoated Protein A/G PLUS-agarose and incubated at 4°C for 16 hr. The immunoprecipitates were isolated by centrifugation and washed five times with IP buffer, and proteins were eluted by heating at 100°C in 62.5 mM Tris-HCl, pH 6.8, containing 2% SDS, 1% β-mercaptoethanol, and 5% glycerol (sample buffer). In some cases, eluted immunoprecipitates were divided into equal halves for immunoblotting analysis.

Immunoblotting

For immunoblotting, SDS-solubilized samples (10 µg) were separated on 8% polyacrylamide gels and transferred to nitrocellulose as described previously (Gurd et al., 1992), and protein blots were reacted with specific antibodies as below. Individual proteins were detected by Western immunoblotting using antibodies against the following: phosphotyrosine (clone 4G10) and c-Src (clone GD11) from Upstate Biotechnology (Lake Placid, NY); NR2A [polyclonal antibody as described by Takagi et al. (1997)], NR2B (clone 13), and Pyk2 (clone 11) from Transduction Laboratories (Lexington, KY); and the phosphoryl group on Tyr1472 of Src family PTKs (Biosource, Camarillo, CA). For some experiments, a rabbit polyclonal antibody specific to phosphorylated Tyr1472 of NR2B was used, which was generously provided by Dr. Tadashi Yamamoto (Department of Oncology, Institute of Medical Science, University of Tokyo, Tokyo, Japan). In some cases, the first antibody was removed by incubation of the blot in 0.1% saline, pH 2.7, at 35°C for 30 min before reaction with a secondary antibody.

Immunoprecipitation and immunoblotting controls

Controls used during blotting runs of IP samples involved either reacting antibodies to lanes loaded only with Protein A/G (to examine nonspecific binding) or to react blots loaded with hippocampal homogenates immunoprecipitated only with preimmune serum. [NMDA-R subunits are robustly expressed in the hippocampus (Petralia et al., 1994a,b)]. An antibody selective for Tyr1472 of the NR2B subunit (Nakazawa et al., 2001) was included in our experiments to independently validate staining obtained with the phosphotyrosine (α-pY) antibody. Also, to examine the specificity of the α-pY antibody, nitrocellulose blots containing proteins from LHA-containing tissue were preincubated with a protein-tyrosine phosphatase (PTPase) to remove the phosphotyrosine residues on all proteins and then incubated with the α-pY antibody. Briefly, the nitrocellulose blots were incubated for 40 min at RT in 2 ml of reaction mixture (as per the supplier’s guidelines; Calbiochem), which contained 50 mM Tris, pH 7.5, 100 mM NaCl, 2 mM EDTA, 5 mM DTT, 0.01% Brij 35, 1 mg/ml BSA, 5 µg/ml each antipain, leupeptin, and aprotinin, 0.1 mM PMSF, and 50 U/ml recombinant Yersinia enterocolitica PTPase (designated Yop51* by Zhang et al. (1992), with a Cys→Arg mutation at residue 235). The nitrocellulose was then washed 3–5 min with TBS containing Triton X-100 and then subjected to immunoblotting as described above.

Samples, detection, visualization, and analysis

Within each treatment group (NMDA- or aCSF-injected in experiment 8a and fasted or ad libitum-fed in experiment 8b), a single sample contained tissue homogenized from LHA-containing punches pooled from at least two rats (i.e., a pool of at least four punches because the punches were bilateral). This was done to ensure that there was enough protein per sample to allow reliable detection. Bound antibodies were detected by enhanced chemoluminescence (SuperSignal; Pierce, Rockford, IL). To obtain optical density (OD) values, exposed X-ray film was scanned using a Bio-Rad (Hercules, CA) GS 700 gel scanner as described previously (Gurd et al., 1992). Because tissue was pooled from subjects within treatment groups as described, each OD score is a quantitation from at least two rats. (For example, a set of five OD scores for a particular band would be derived from tissue samples harvested from a total of at least 10 rats undergoing the same experimental manipulation.) Once OD values were obtained, a mean OD value ± SEM was calculated for the group. Mean scores were analyzed by an unpaired t test or for, nonparametric analysis, by a Mann–Whitney rank sum test, with α set at 0.05.

Results

Genistein suppressed NMDA-elicted eating in a behaviorally selective manner (experiments 1–3)

Figure 1A shows the results of experiment 1a. As previously reported (Stanley et al., 1993b,c, 1996, 1997; Khan et al., 1999), a unilateral LHA microinjection of NMDA, with a Cys→Arg mutation at position 850, caused substantial food intake at all times (7.9 ± 1.3 gm after NMDA compared with 0.6 ± 1.4 gm after vehicle at 0.5 hr; p < 0.05 by one-way ANOVAs). A new finding was that pretreatment with LHA injection of the PTK inhibitor genistein (chemical structure, Fig. 1B) markedly reduced NMDA-elicted eating, with a maximal suppression of 53% to 3.7 ± 1.3 gm at 0.5 hr after injection. Two-way ANOVA of the genistein and NMDA versus the vehicle and NMDA treatments revealed a significant effect of treatment (F1,172 = 8.9; p < 0.01) but no effect of time after injection or their interaction.
food intake ($F_{(3,204)} = 13.3; p < 0.0001$), with no significant interaction. In contrast, multiple comparisons revealed no significant effect of genistein at any time after injection. Visual inspection suggests that the slight overall effect was primarily attributable to a delayed suppression (i.e., at 4 hr) produced by the 2.4 nmol dose of genistein. Most importantly, the 240 nmol dose of genistein, which produced 53 and 62% suppressions of NMDA-elicited eating 30 min after injection in experiments 1 and 2, respectively, had no effect on KA-elicited eating at that time.

Finally, in contrast to its effects on NMDA-elicited feeding, genistein did not suppress deprivation-elicited eating (experiment 3) at any postinjection time measured (e.g., at 0.5 hr after injection, vehicle- and genistein-injected rats ate 17.1 ± 1.4 and 16.5 ± 1.6 gm, respectively; at 4 hr, they ate 21.7 ± 1.4 and 22.1 ± 1.5 gm, respectively). That genistein did not suppress KA- or deprivation-elicited eating argues that its suppressive effects on NMDA-elicited eating were not caused by the induction of nausea or malaise because rats injected with genistein were still capable of robust eating responses and, in the case of KA-elicited eating, especially during the time when absolute KA-elicited intake was the largest (i.e., the first 30 min). Critically, the data also suggest that PTKs may not be essential in all LHA mechanisms of eating stimulation. They further suggest that the mechanisms underlying eating attributable to NMDA-R and KA-R activation in the LHA are distinct, a conclusion supported by a recent study of KA-R-delimited signaling (Rozas et al., 2003).

Tyrophostin A48 suppressed NMDA-elicited feeding (experiment 4)

Although they can affect at least one non-PTK target (Martin, 1998), the tyrophostins are reportedly more selective than genistein at inhibiting PTKs because they act at the substrate-binding site in addition to or instead of the ATP-binding site of the PTK (Levitzki, 1992). Tyrophostin A48 (chemical structure, Fig. 3A), also suppressed NMDA-elicited feeding ($F_{(3,132)} = 3.0; p < 0.05$ by two-way ANOVA excluding vehicle), although the effect was weaker than that produced by genistein. Multiple comparisons revealed that only the highest dose of Tyr A48 (212 nmol) significantly suppressed NMDA-elicited food intake (e.g., from 10.5 ± 1.9 to 6.9 ± 1.8 gm at 0.5 hr after injection) (Fig. 3A).

PP1, an Src family-selective PTK inhibitor, robustly suppressed NMDA-elicited feeding (experiment 5)

PP1 (chemical structure, Fig. 3B) is a selective pyrazolopyrimidine class of PTK inhibitor that acts as an ATP analog. It is orders of magnitude more potent of an inhibitor of Src family PTKs than of other PTK targets ($IC_{50}$ values, 0.005–0.17 vs 50–100 μM, respectively) and only minimally affects other kinase targets (Hanke et al., 1996; Liu et al., 1999; Waltenberger et al., 1999; Susa et al., 2000). Additionally, PP1 is two orders of magnitude more potent than genistein in inhibiting Src PTKs (cf. Akiyama and Ogawara, 1991; Hanke et al., 1996; Waltenberger et al., 1999; Susa et al., 2000). In experiment 5a, as shown in Figure 3B, LHA-administered NMDA again elicited significant food intake at all
postinjection times measured (e.g., at 0.5 hr, NMDA elicited 6.8 ± 1.9 gm of food intake versus 0.2 ± 0.1 gm for vehicle; p < 0.05 by one-way ANOVAs for each time point). A new finding is that PP1 produced a strong dose-dependent suppression of the NMDA-elicited response (Fig. 3B), with 0.2, 2.1, and 21.2 nmol of PP1 suppressing NMDA-elicited feeding by 37% (4.3 ± 2.3 gm), 54% (3.1 ± 1.4 gm), and 69% (2.2 ± 1.2 gm), respectively (F(2,135) = 4.6; p < 0.01 by two-way ANOVA excluding vehicle- and inhibitor-only treatments; p < 0.05 by the Student–Neuman–Keuls method).

In experiment 5b (data not shown), the suppression of NMDA-elicited eating by PP1 was reproduced in a naive group of animals (10.5 ± 2.3 to 6.2 ± 1.6 gm at 0.5 hr; p < 0.05 by the Student–Neuman–Keuls method).

PP1 suppressed feeding triggered by food deprivation (experiment 6)

To help determine the contribution of Src family PTKs to feeding behavior caused by food deprivation, we bilaterally injected PP1 into the LHA of rats deprived of food for ~24 hr. As shown in Figure 4A, vehicle-injected rats ate 13.8 ± 1.8 gm at 0.5 hr after injection, a score that was suppressed by ~24% in PP1-injected animals (10.5 ± 2.4 gm). A two-way ANOVA revealed significant effects of both treatment (F(1,68) = 6.7; p < 0.05) and time after injection (F(3,68) = 7.1; p < 0.001) on food intake scores but not their interaction (p = 0.88).

PTK inhibitors differed in their power and potency to suppress NMDA-elicited eating

We compared the efficacy of all three PTK inhibitors to suppress NMDA-elicited eating behavior across the three sets of studies. Figure 4B shows that PP1 produced the largest suppression of NMDA-elicited feeding (69% at the highest dose tested), followed by genistein (53% at the highest dose tested) and Tyr A48 (34%). PP1 was also the most potent inhibitor. Although it was three orders of magnitude lower in concentration, the lowest effective dose of PP1 produced a suppression similar in magnitude to that produced by the highest dose of Tyr A48 (Fig. 4B).

Src-like immunoreactivity was localized within lateral hypothalamic neurons (experiment 7)

To determine whether Src PTK exists in LHA cells, we performed immunohistochemistry using a polyclonal antibody that recognizes Src. Figure 5A shows that subpopulations of LHA neurons displaying robust Src-like immunoreactivity were present in an area ranging from the perifornical area to the margin of the optic tract. Immunoreactive cells were identified as neurons on the basis of their similarity to LHA neurons described in a Golgi study (Millhouse, 1979). Staining was observed in both somata and neurites (Fig. 5B), and high-power visualization under oil immersion revealed that some of these neurites were probably dendrites, as multiple neurites were often seen extending from a single cell body. Staining was also observed in the nuclei of these cells, which was absent in control sections processed without primary antibody, suggesting that Src-like immunoreactivity is detectable in multiple cellular compartments. Our results support and supplement those of Ross et al. (1988) and Sugrue et al. (1990), who provided brief textual and photographic descriptions, respectively, of Src isoforms in the LHA as part of larger anatomical surveys.

In contrast to the robust immunostaining we observed using this antibody, control sections processed in its absence displayed virtually no immunoreactivity (data not shown). Moreover, because the “background staining” observed in the sections treated with the primary antibody was eliminated in the control sections, some of this staining might be from cut immunoreactive neurites out of the plane of section of the tissue.

Food deprivation but not LHA NMDA injection was associated with altered protein-tyrosine phosphorylation of several proteins in LHA-containing tissue

Because some PTK inhibitors were able to suppress feeding triggered not only by NMDA injection but also by food deprivation, we next determined the effects of these feeding conditions on protein-tyrosine phosphorylation in LHA-containing tissue. In experiment 8a, micropunched LHA-containing tissue was removed from rats receiving bilateral injections of aCSF vehicle or NMDA. Figure 6A shows that prominent tyrosine-phosphorylated (Tyr-P) protein components were observed in the 50–200 kDa molecular weight range, including bands of ~60, 95, 115, and 180–190 kDa. We focused our quantitative analysis on the p60 and p180–190 bands because Src family PTKs are detectable in multiple cellular compartments. Our results supported and supplemented those of Ross et al. (1988) and Sugrue et al. (1990), who provided brief textual and photographic descriptions, respectively, of Src isoforms in the LHA as part of larger anatomical surveys.

In contrast to the robust immunostaining we observed using this antibody, control sections processed in its absence displayed virtually no immunoreactivity (data not shown). Moreover, because the “background staining” observed in the sections treated with the primary antibody was eliminated in the control sections, some of this staining might be from cut immunoreactive neurites out of the plane of section of the tissue.
Food deprivation but not LHA NMDA injection was associated with altered phosphorylation of Src and altered expression of Pyk2

We next examined whether Src and Pyk2 PTK expression and the phosphorylation of Src family PTKs at Tyr\(^{527}\) differed between aCSF and NMDA or fed and fasted groups. The phosphorylation of Src family PTKs at Tyr\(^{527}\) was examined because depolymerization of this regulatory site is a well-known mechanism that relieves Src family PTKs from their autoinhibitory confirmation and facilitates their increased activation (Cooper et al., 1986; Thomas and Brugge, 1997; Xu et al., 1999). Thus, the phosphorylation state of this site provides one important index of apparent catalytic activity (Fig. 7C) (for details, see Discussion). Figure 7A shows that no statistically significant differences were observed in the phosphorylation levels of Src family PTKs at Tyr\(^{527}\), total Src protein, or total Pyk2 protein between aCSF- and NMDA-injected rats. Although such changes could have occurred in a subset of LHA neurons but that remained undetectable within the larger micropuncched samples we assayed, these results suggest that NMDA injection is not sufficient to drive such changes in vivo. We cannot, however, rule out mechanisms that activate Src PTKs independent of its dephosphorylation at Tyr\(^{527}\) (Ma et al., 2000) or the possibility that the anesthesia might have masked effects that would otherwise have been apparent.

In experiment 8b, identical measures were made on LHA-containing tissue rats fasted for 48 hr or their ad libitum-fed controls. Figure 7B shows that, in contrast to NMDA injection, food deprivation significantly changed two of these three end points. Specifically, there was a marked decrease in the mean OD value for immunoreactive pSrc\(^{527}\) in LHA-containing tissue of the fasted rats relative to controls (0.3 ± 0.1 compared with 0.8 ± 0.1, respectively; \(p < 0.01\) by t test) (Fig. 7B). However, the total expression of Src remained unchanged between groups (Fig. 7B). Also, levels of the Src-activating PTK Pyk2 were robustly elevated in LHA-containing tissue from fasted rats (Fig. 7B) (1.4 ± 0.2 compared with 0.6 ± 0.1, respectively; \(p < 0.05\) by t test). We examined Pyk2 because of recent studies demonstrating it is a critical participant within signaling pathways that use NMDA-R activation and Src activity (Huang et al., 2001).

Tyrosine phosphorylation of LHA NMDA-R subunits did not change in association with NMDA injection or fasting

PTKs such as Src can enhance NMDA-R activity (Wang and Salter, 1994; Chen and Leonard, 1996; Köhr and Seeberg, 1996; Yu et al., 1997; Lancaster and Rogers, 1998; Xiao et al., 2000). This enhancement may occur as a result of NMDA-R subunit phosphorylation by Src, although this awaits empirical demonstration (Salter and Kalia, 2004). NR2A and NR2B subunits, which can be tyrosine-phosphorylated (Moon et al., 1994; Lau and Huganir, 1995), might also be within LHA NMDA-Rs active during feeding stimulation (Petralia et al., 1994a,b; Khan et al., 1999, 2000b).

Accordingly, we hypothesized that, by phosphorylating these subunits in LHA neurons, PTKs could contribute to changes in

Other bands in the 75–110 kDa range from fasted rats also displayed apparent changes in optical density; these, however, were not quantitatively analyzed. In contrast, there were no significant differences in the Tyr-P levels of p180–190 between the two groups.

To verify the specificity of our phosphotyrosine antibody, we performed a control experiment in which we first exposed the immunoblot to a PTPase and then reacted it with the phosphotyrosine antibody. As shown in Figure 6C, pretreating the blot with PTPase eliminated phosphotyrosine immunostaining of all separated proteins.
food intake and body weight by enhancing LHA NMDA-R activity (Khan et al., 1999) (A. M. Khan, E. R. Gillard, B. G. Stanley, Phosphorylation of Hypothalamic NMDA Receptors as a Possible Mechanism Contributing to Changes in Food Intake and Body Weight, in Matters Arising, International Society for Neurochemistry, available on-line). In such a case, feeding stimulation should increase the Tyr-P of LHA NR2A and NR2B subunits.

However, we found that NMDA injection did not appreciably change the phosphorylation of NR2B at Tyr1472 (data not shown). Additionally, Figure 8 shows that fasting did not alter levels of protein or Tyr-P for NR2A and NR2B subunits immunoprecipitated from LHA-containing tissue (mean OD scores, 2.09 ± 0.64 and 1.69 ± 0.82, respectively; p = 0.54 by Mann–Whitney rank sum test). Nor did fasting significantly enhance Tyr-P of NR2B at Tyr1472 (data not shown) (mean OD scores, 2.91 ± 0.51 and 2.67 ± 0.79; p = 0.80 by t test). Thus, neither NMDA injection nor food deprivation appears to be associated with changes in the Tyr-P levels of the NR2A and NR2B NMDA-R subunits, although such changes may occur in only subpopulations of LHA neurons and, therefore, at levels that escaped detection by our methods. However, these results do confirm reports of NR2A and NR2B NMDA-R subunits in the LHA (Khan et al., 1999, 2000b) and also provide the first evidence that these subunits are tyrosine-phosphorylated in this region.

**Discussion**

The present study provides four lines of evidence suggesting that Src family PTKs and Tyr-P participate in LHA signaling mechanisms underling feeding stimulation: (1) NMDA-elicited feeding is powerfully suppressed by structurally distinct LHA-administered PTK inhibitors, including one selective for Src family PTKs; (2) deprivation-induced feeding is also significantly suppressed by the Src family-selective inhibitor; (3) Src is robustly expressed in LHA neurons and, along with Pyk2 PTK and various Tyr-P proteins, is detectable in LHA-containing tissue; and (4) food deprivation is associated with altered Tyr-P of various LHA proteins, an increase in the apparent activity of one or more Src family PTKs, and a robust increase in Pyk2 expression.

PTK inhibitors suppress eating

LHA administration of the PTK inhibitor genistein marked suppression NMDA-elicited eating. Because genistein can affect non-PTK targets, we evaluated its chemical and behavioral specificity for producing these effects. First, daidzein, a genistein analog that does not inhibit PTKs, did not suppress NMDA-elicited eating, suggesting that the effects of genistein are specific to its PTK inhibitory actions.

Figure 6. Tyrosine-phosphorylated proteins are present within total protein extracts of homogenized LHA-containing tissue. A, B, Total protein within LHA-containing tissue from aCSF- and NMDA-injected rats (A) or ad libitum-fed or 48 hr-fasted rats (B) was probed with an α-pY antibody. Positions of molecular weight standards are along the left of the blots (in kilodaltons). Note the prominent —60, —95, and —180–190 kDa bands. Data are representative of four experiments, each performed using tissue obtained from one or two animals. D, E, Tyr-P levels are expressed as OD units ± SEM for the 60 and 180–190 kDa bands in tissue from aCSF- and NMDA-injected rats (D) or ad libitum-fed and 48 hr-fasted rats (E). Compared with controls, fasted rats had significantly decreased Tyr-P for the 60 kDa band (inverted carat; p < 0.05). C, Specificity of the α-pY antibody. The α-pY immunoreactivity for total protein from LHA-containing tissue was abolished with (+) but not without (−) tyrosine phosphatase pretreatment.
across similar but not identical assay conditions, apparently inhibits Src family PTKs with a potency at least two orders of magnitude greater than that for genistein (Susa et al. 2000). Although our data do not eliminate the possibility of effects on other targets, these facts, along with the observation that PP1 produced the most potent and powerful suppression of NMDA-elicited feeding, strongly implicate Src family PTKs as the neural substrates most sensitive to signals triggering such feeding. In support of this, PP1 but not genistein suppressed deprivation-elicited feeding, underscoring that perhaps Src PTKs are involved. Thus, our pharmacological studies suggest that PTKs, particularly Src family PTKs, are involved in mechanisms underlying feeding stimulation.

Src, Pyk2, and various other tyrosine-phosphorylated proteins exist in the LHA. Our quantitative immunochemical data also support a role for PTK signaling within LHA neurons, revealing robust expression of Src, Pyk2, and several as yet unidentified tyrosine-phosphorylated proteins in LHA-containing tissue. Our Pyk2 detection confirms, at the protein level, studies localizing transcripts for Pyk2 variants in the hypothalamus (Xiong et al., 1998; Menegon et al., 1999). Also, Src was immunolocalized to selected perifornical LHA neurons, which contain neural sub-
strates sensitive to the feeding stimulation effects of neuropeptide Y, hypocretin/orexin, or Glu (Stanley et al., 1992, 1993a,b,c; Khan et al., 2000b; Marcus et al., 2001; Backberg et al., 2002; Lee and Stanley, 2002; Wolak et al., 2003), suggesting that Src can interact with signaling components of these neural substrates to influence feeding or other behaviors.

Effects of NMDA injection or food deprivation on LHA tyrosine phosphorylation, PTK activity, and PTK expression

The first major finding from the biochemistry was that deprivation-induced feeding but not NMDA-elicited feeding was associated with a marked decrease in the Tyr-P of the p60 band. Second, because Src family PTKs (EC 2.7.1.112) are themselves regulated by Tyr-P and are ~60 kDa (Bjorge et al., 2000; Tatosyan and Mizzenina, 2000), we asked whether the decreased Tyr-P of p60 might reflect decreased Tyr-P of Src family PTKs. Accordingly, we reacted immunobLOTS of micropunched LHA-containing tissue with a phospho-specific Src family PTK antibody directed against Tyr-527, located near the C-terminus. Tyr-P of this regulatory site inhibits Src catalytic activity and is a well-studied mechanism of Src regulation (Fig. 7C) (Cooper et al., 1986; Sancha and Kuriyan, 1997; Hubbard and Till, 2000; Wang et al., 2001; Young et al., 2001). This antibody revealed markedly reduced Tyr-P at this site in food-deprived rats, suggesting that food deprivation markedly desinhibits and perhaps activates one or more Src family PTKs. This suggestion is supported by the suppression of deprivation feeding we observed using the Src family-selective PTK inhibitor PP1. Food-deprived rats also displayed elevated expression of Pyk2, a soluble PTK known to interact with Src in some neural systems (Ali and Salter, 2001; Huang et al., 2001; Salter and Kalia, 2004). In contrast, NMDA-elicited feeding stimulation was not associated with elevated Src family activity or Pyk2 expression. Collectively, these results strongly implicate Src family PTKs in LHA signaling mechanisms underlying feeding stimulation produced by food deprivation.

Interpreting the differential effects of NMDA-elicited and deprivation-induced feeding stimulation on PTK activity and expression

Our behavioral data demonstrate that LHA injections of PTK inhibitors, including one selective for the Src family, are more effective in suppressing NMDA-elicited feeding than food deprivation-induced feeding. In contrast, the biochemical data reveal that food deprivation is more effective than NMDA injection in driving detectable changes in the activity and expression of PTKs in LHA-containing tissue. Thus, food deprivation is sufficient to drive changes in PTK activity, but PTKs are not so critical for deprivation feeding to occur. Conversely, NMDA-elicited feeding stimulation is not sufficient to increase PTK activity, but PTKs appear to be quite critical for the full magnitude of the NMDA-elicited feeding response.

These biochemical and behavioral data together form the basis of the model depicted in Figure 9. The model posits that deprivation feeding will activate more distinct types of neural substrate than feeding driven by LHA NMDA injection (see Orexigenic Signals in both panels of Fig. 9). This model is an elaboration of the prevailing view that food deprivation engages multiple neural circuits and chemical messengers (Berthoud, 2002; Saper et al., 2002) and adds to that formulation the suggestion that food deprivation recruits a greater variety of signaling pathways to elicit feeding than does acute NMDA injection. From this, the model predicts that food deprivation, by engaging many signaling pathways to elicit feeding, would be suppressed to only a minor extent with PTK inhibitors but would still markedly shift the activity and expression of PTKs in the LHA (Fig. 9, right panel). In contrast, although acute NMDA injection triggers feeding behavior that is more easily suppressed by PTK inhibitors and that, therefore, may rely more heavily on PTK signaling mechanisms, such stimulation is not sufficient to alter PTK activity or expression as measured by the indices we used. This is perhaps because the NMDA-R was the only signal, of many possible PTK-linked signals, that was engaged by our NMDA injection (Fig. 9, compare left, right panels).

Using in vivo approaches to identify intracellular effectors is challenging but is clearly important for understanding their function in the context of behavior (Dumont et al., 2002). The present work supports the argument that such an understanding is more likely to be achieved in the intact animal by using multiple stimulation paradigms and by using a combination of biochemistry and behavioral analysis to help sharpen and constrain the interpretations that might otherwise be drawn when using only one of these approaches.
Src family PTKs in LHA neurons may integrate feeding signals

Src family PTKs can be activated by G-protein-coupled receptors (GPCRs) or receptors coupled to Tyr-P mechanisms (Ma et al., 2000; Ram and Iyengar, 2001); activation of either receptor class could mobilize Src family PTKs in LHA neurons during food deprivation. For example, LHA afferents from the arcuate nucleus (Elias et al., 1998) could release NPY, AgRP, and α-MSH to activate GPCRs. In contrast, Tyr-P-linked activation of a leptin receptor isoform (Li and Friedman, 1999; Hegyi et al., 2004) described in the LHA (Fei et al., 1997; Elmquist et al., 1998; Iqbal et al., 2001) could trigger downstream effectors that potentially interact with Src (Chaturvedi et al., 1998; Hung and Elliott, 2001). Additionally, serine-threonine protein kinases implicated in feeding stimulation (Gillard et al., 1997, 1998a,b; Sheriff et al., 1997) could cross-signal with LHA PTKs (Nijhoff et al., 2000; Schmitt and Stork, 2002). How Src family PTKs help transfer feeding signals within LHA neurons is unclear, but our results suggest that some of this transfer involves Pyk2 and Src recruitment without significant alterations in NMDA-R subunit Tyr-P.

Concluding remarks

These studies constitute the first description of PTK involvement in LHA mechanisms underlying feeding stimulation. Specifically, Src family PTKs, which are robustly expressed within LHA neurons, are apparently activated after food deprivation and appear critical for ensuring the full magnitude of food intake that occurs during acute and prolonged feeding stimulation. That soluble PTKs such as Src mediate signals to help control vital behaviors such as eating underscores the importance of these enzymes for the survival of multicellular organisms, which, in contrast to their unicellular relatives, may be unique in having these versatile proteins encoded within their genomes (Hanks and Hunter, 1995; Darnell, 1997).

References

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