Transduction in Olfactory Receptor Neurons of *Xenopus laevis* Larvae: Pharmacological Blockage with FM1-43 and Endocannabinoid Modulation

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submitted by
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Göttingen, September 7th, 2009

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Für meine Eltern
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<table>
<thead>
<tr>
<th>Abbreviation</th>
<th>Description</th>
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<tbody>
<tr>
<td>2-AG</td>
<td>2-arachidonoylglycerol</td>
</tr>
<tr>
<td>ATP</td>
<td>Adenosine triphosphate</td>
</tr>
<tr>
<td>[Ca^{2+}]_i</td>
<td>Intracellular Ca^{2+} concentration</td>
</tr>
<tr>
<td>cAMP</td>
<td>Cyclic adenosine monophosphate</td>
</tr>
<tr>
<td>CB</td>
<td>Cannabinoid</td>
</tr>
<tr>
<td>CE</td>
<td>SE(5-(and-6)-carboxyeosin diacetate, succinimidyl ester)</td>
</tr>
<tr>
<td>cGMP</td>
<td>Cyclic guanosine monophosphate</td>
</tr>
<tr>
<td>CNG</td>
<td>Cyclic nucleotide-gated</td>
</tr>
<tr>
<td>DAGLα, β</td>
<td>Diacylglycerol lipase α, β</td>
</tr>
<tr>
<td>DMSO</td>
<td>Dimethyl sulfoxide</td>
</tr>
<tr>
<td>EDTA</td>
<td>Ethylenediaminetetraacetic acid</td>
</tr>
<tr>
<td>EGTA</td>
<td>Ethylene glycol tetraacetic acid</td>
</tr>
<tr>
<td>FAAH</td>
<td>Fatty acid amide hydrolase</td>
</tr>
<tr>
<td>FM1-43</td>
<td>N-(3-triethylammoniumpropyl)-4-(4-(dibutylamino)styryl)pyridinium dibromide</td>
</tr>
<tr>
<td>G_{olf}</td>
<td>Olfactory-specific guanosine triphosphate (GTP)-binding protein</td>
</tr>
<tr>
<td>GTP</td>
<td>Guanosine triphosphate</td>
</tr>
<tr>
<td>HEPES</td>
<td>4-(2-hydroxyethyl)-1-piperazineethanesulfonic acid</td>
</tr>
<tr>
<td>MAGL</td>
<td>Monoacylglycerol lipase</td>
</tr>
<tr>
<td>MAP</td>
<td>Mitogen-activated protein</td>
</tr>
<tr>
<td>NAPE-PLD</td>
<td>N-acylphosphatidylethanolamide-hydrolyzing phospholipase D</td>
</tr>
<tr>
<td>NCX</td>
<td>Sodium calcium exchanger</td>
</tr>
<tr>
<td>NMDG</td>
<td>N-Methyl-D-glucamine</td>
</tr>
<tr>
<td>OE</td>
<td>Olfactory epithelium</td>
</tr>
<tr>
<td>ORN</td>
<td>Olfactory receptor neuron</td>
</tr>
<tr>
<td>PCR</td>
<td>Polymerase chain reaction</td>
</tr>
<tr>
<td>PMCA</td>
<td>Plasma membrane calcium ATPase</td>
</tr>
<tr>
<td>RT PCR</td>
<td>Reverse transcription polymerase chain reaction</td>
</tr>
<tr>
<td>TAE</td>
<td>Tris acetate EDTA</td>
</tr>
</tbody>
</table>

Abbreviations exclusively used in figures are explained in the respective figure legends.
“Smell is a potent wizard that transports us across thousands of miles and all the years we have lived. The odors of fruits waft me to my southern home, to my childhood frolics in the peach orchard. Other odors, instantaneous and fleeting, cause my heart to dilate joyously or contract with remembered grief. Even as I think of smells, my nose is full of scents that start awake sweet memories of summers gone and ripening fields far away.”

Helen Keller

*US blind & deaf educator (1880-1968)*
1 Introduction

The sense of smell enables almost all species of the animal kingdom to recognize and discriminate between a large array of molecules called odors with a great accuracy and sensitivity. Especially in humans, this sense was regarded as a kind of luxury, but in fact, animals heavily depend on detecting molecules in the environment in order to survive as a successful organism. This makes olfaction one of the most remarkable senses, but it is also the least understood.

The olfactory system is a chemosensory system, indispensable for the interplay of an organism with its environment. This sense is of particular importance as it allows to identify about 10,000 odors (for humans, Mombaerts, 2004), and it has a tremendous impact on a variety of behavioral patterns. Olfaction is a significant sensory input for appetite regulation and food seeking behavior. Most animals including humans use olfactory information in order to appreciate food palatability and to initiate food intake (Rolls, 2005; Yeomans, 2006). Furthermore, the olfactory system influences emotional responses like anxiety, fear, and pleasure, reproductive functions like sexual and maternal behaviors as well as social behaviors like recognition of conspecifics and predators (Schultz and Tapp, 1973; Nimmermark, 2004; Takahashi et al., 2005). Olfaction is often underestimated, but its significance becomes obvious in the case of its loss. Many patients with impaired olfaction have a poor quality of life and develop feelings of personal isolation, lack of interest in eating, and emotional blunting. Moreover, olfactory disorders are associated with depression (Toller, 1999; Nordin and Brämerson, 2008). It is now apparent that dysfunction of olfaction is one of the first symptoms of neurodegenerative diseases like Alzheimer’s and Parkinson’s disease (Doty, 2008; Doty, 2009), and it is associated with many other cognitive diseases like schizophrenia or Huntington’s disease (Lombion-Pouthier et al., 2006; Lazic et al., 2007; Atanasova et al., 2008).

The increasing understanding of the links between olfaction and various diseases will eventually lead to the discovery of new disease mechanisms, which in turn introduce new targets for drug development. This is of particular importance since none of the diseases mentioned above can be cured to date. However, before functional links in this field can be established, further knowledge about olfaction
itself has to be gained. Regarding the first steps of peripheral odor encoding, two questions need to be addressed for a better understanding of the mechanisms underlying olfaction:

*By which mechanisms are odors coded in the neuronal substrates?*

*How is odor coding modulated by intrinsic and extrinsic factors?*

In this thesis, significant findings are presented that contribute to answer these questions. Before showing the results I obtained during the last three years, I would like to introduce basic principles of olfaction, as far as they are important for a better understanding of the results. Section 1.1 summarizes the organization of the olfactory mucosa and principles of odor encoding by olfactory receptor neurons. In section 1.2 the current understanding of modulatory principles in the olfactory mucosa are summarized. Especially the endocannabinoid system is highlighted: on the one hand the importance of this system for olfaction is elaborated, on the other hand the common physiological importance of endocannabinoids and their mechanism of action are illustrated.

### 1.1 The sense of smell

#### 1.1.1 Morphology of the olfactory system

The olfactory system in vertebrates can roughly be divided into the olfactory epithelium (OE), the olfactory bulb, and higher brain centers. The first steps of odor detection take place in the OE, whereas further processing takes place centrally. The OE is located in the nasal cavity embedded under a layer of mucus, which is secreted by olfactory glands (primarily Bowman’s glands) and sustentacular cells (Getchell, 1986; Gold, 1999, Schwob, 2002). This epithelium contains three main cell types (Figure 1):

1. Olfactory receptor neurons (ORNs; Figure 1, upper part, orange) are bipolar neurons. These primary sensory cells transduce and transform the binding of ligands to olfactory receptors into sequences of action potentials. From the
1.1 The sense of smell

small soma, a single dendrite extends to the nasal cavity and ends in a knob bearing cilia. In cilia olfactory transduction takes place upon binding of odors to the receptors situated in the membrane. On the basal side of the soma an unmyelinated axon originates and projects via the olfactory nerve to the olfactory bulb (Getchell, 1986; Schild and Restrepo, 1998; Gold, 1999).

2. Sustentacular cells (Figure 1, purple) have multi-faceted functions in the OE. Besides their role in mucus secretion and regulation of the ionic content of the mucus (Getchell and Getchell, 1992; Hansen et al., 1998) they have a glia-like function and insulate ORNs physically and chemically (Breipohl et al., 1974; Getchell and Getchell, 1992), phagocytose dead cells (Suzuki et al., 1996),

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**Figure 1: Organization of the olfactory system.** The morphology of the olfactory system is illustrated in this sketch. Figure modified from Lang and Lang (2007); In “Basiswissen Physiologie”, page 429; 2nd edition; Springer Berlin Heidelberg Verlag.
and detoxify noxious substances (Lazard et al., 1991). Recently it was suggested that sustentacular cells play a role in intraepithelial signaling, e.g. in cell proliferation by purinergic signaling (Hegg and Lucero, 2006; Hassenklöver et al., 2008; Hassenklöver et al., 2009). Sustentacular cells are arranged columnar at the apical side of the OE and are either ciliated or secretory.

3. Basal cells (Figure 1, upper part, green) represent the stem cells within the OE and are responsible for cell regeneration. The life span of ORNs is 30 to 60 days and they have to be replaced continuously due to the exposure of the OE to a variety of toxic agents. The consequential cell damage and death is compensated by differentiation of basal cells into ORNs and sustentacular cells (Schwob, 2002; Ronnett and Moon, 2002).

Each mature ORN relays electric information via its axon running in the olfactory nerve to the olfactory bulb, which is separated from the OE by the cribiform plate in adult vertebrates (Figure 1). There, it forms synapses in specialized structures called glomeruli. Two types of interneurons are involved in the olfactory processing in the bulb: periglomerular cells and granule cells. The output neurons of the olfactory bulb, the mitral cells, are glutamatergic and convey olfactory information to the primary olfactory cortex (Mori et al., 1999; Lledo et al., 2005; Wilson and Mainen, 2006).

1.1.2 Olfactory transduction and transformation in ORNs

Olfactory transduction
The first step in perceiving an odor is the interaction of the odorant molecule with the olfactory receptor on cilia of ORNs (Figure 2). These receptors belong to the family of seven transmembrane receptors and are encoded by about 1000 separate genes in rat and mouse and 400 in frog (Buck and Axel, 1991; Mombaerts, 1999; Niimura and Nei, 2006). The interaction of odors with the receptors leads to the intracellular activation of an olfactory-specific guanosine triphosphate (GTP)-binding protein called $G_{olf}$. The $\alpha$-subunit of this G protein in turn mediates the activation of adenylate
1.1 The sense of smell

The sense of smell involves the binding of an odor molecule to olfactory receptors in the ciliary membrane, leading to a signal transduction cascade. This process is primarily mediated by adenylate cyclase III, a membrane-bound enzyme that catalyzes the reaction of ATP to cAMP. This reaction opens cyclic nucleotide-gated (CNG) channels that are permeable for cations, including Na\(^+\) and Ca\(^{2+}\) (Dhallan et al., 1990). The subsequent inward flow of Na\(^+\) and Ca\(^{2+}\) depolarizes the cell membrane and opens Ca\(^{2+}\)-activated Cl\(^-\) channels (Stephan et al., 2009), resulting in an efflux of Cl\(^-\) ions. This leads to a further depolarization due to the elevated equilibrium potential of Cl\(^-\). The cell membrane is thus further depolarized (Schild and Restrepo, 1998; Kleene, 2008).

The cAMP-dependent transduction mechanism appears to be predominant particularly in terrestrial vertebrates. However, besides this well described pathway also non-cAMP-dependent transduction mechanisms exist (Schild and Restrepo, 1998; Manzini et al., 2002). These comprise inositol 1,4,5,-tris-polyphosphate as a second messenger (Kaur et al., 2001; Bruch, 1996) as well as nitric oxide and carbon

---

**Figure 2: cAMP-dependent olfactory transduction.** Upon binding of an odor (green square) to an olfactory receptor (R) in the ciliary membrane, GTP bound to G\(_{olf}\) is hydrolyzed. This leads to the dissociation of the α-subunit which activates the adenylate cyclase (AC). Subsequently, cAMP gates CNG channels which are permeable for cations. Influx of Na\(^+\) and Ca\(^{2+}\) leads to both depolarization of the membrane and opening of Ca\(^{2+}\)-dependent Cl\(^-\) channels which results in a further depolarization by a Cl\(^-\) efflux. Figure modified from Firestein (2001).
monoxide (Breer and Shepherd, 1993; Broillet and Firestein, 1996). Furthermore, odors induce not only excitatory responses in ORNs, but they can also elicit inhibitory responses in other ORNs (Vogler and Schild, 1999; Morales et al., 1997).

**Olfactory transformation**

The receptor potential, which is generated in cilia, propagates electrotonically to the soma. Because of their small membrane capacitance, high membrane resistance, and long time constant (Schild et al., 1994; Schild and Restrepo, 1998; Imanaka and Takeuchi, 2001) ORNs are highly sensitive to odorant stimulation. If the potential at the soma reaches threshold, voltage-gated Na$^+$ channels open at the axon hillock, thus generating an action potential. This strong depolarization then activates voltage-gated Ca$^{2+}$ channels (high-voltage-activated, activated between -30 mV and -40 mV) at the proximal dendrite. Ca$^+$-dependent K$^+$ channels at the proximal dendrite and voltage-gated K$^+$ channels at the axon hillock serve to repolarize the cell membrane and to terminate the action potential (Schild, 1989; Schild et al., 1994). Action potentials are conveyed via the olfactory nerves towards the olfactory bulb where further processing takes place (Wilson and Mainen, 2006; Toida, 2008).

1.1.3 *Peripheral odor coding*

Olfactory receptor neurons detect many odors with their qualitative, quantitative, and temporal information. A single odorant can activate multiple olfactory receptors and one olfactory receptor can bind many odorants. The hypothesis of one olfactory receptor type per ORN is reevaluated and the probability of the expression of more than one olfactory receptor per ORN is discussed (Rawson et al., 2000; Mombaerts, 2004; Tian and Ma; 2008).

Moreover, the OE consists of ORN subsets, which can be classified e.g. according to their odor sensitivity, maturation state, and transduction cascade. Regarding the second messenger cascades, there exist multiple types in the OE of various species (Schild and Restrepo, 1998; Manzini et al., 2002) and some may even be coexpressed within one cell (Ko and Park, 2006). This means that odor
coding is a complex process and that ORN subsets work not only in parallel but may also interact or are coactivated.

In order to understand peripheral odor coding mechanisms, ORN subsets need to be characterized in detail. Tools, allowing interference with single ORN subtypes are required. To date, the differentiation of ORN subtypes according to the olfactory receptor expression is possible by genetic manipulation (Mombaerts et al., 1996; Grosmaître et al., 2005). Differentiation of ORNs according to their sensitivity is mainly possible by stimulating ORNs with odors or second messenger analogues (Manzini and Schild, 2003a; Gautam et al., 2006).

Having a closer look at the cAMP-dependent ORN subset, a well known feature of olfactory transduction is the expression of a cascade of two generator channels, a Ca\(^{2+}\)-permeable CNG channel driving a Ca\(^{2+}\)-dependent Cl\(^-\) channel (Stephan et al., 2009). The transduction of odorants can be interfered with on three levels: First, at the level of olfactory receptors (Oka et al., 2004a; Oka et al., 2004b; Sanz et al., 2005), second, at the level of receptor potential modulation (e.g. Czesnik et al., 2007 (cannabinoids); Bouvet et al., 1988 (acetylcholine); Kawai et al., 1999 (adrenaline)), and third, at the level of action potential generation. Blocking olfactory transduction at the level of one or the other the generator channel has proven difficult so far, due to the lack of specific Cl\(^-\) channel blockers and the lack of CNG channel blockers that act at resting membrane potentials.

Obviously, such blockers would be particularly important in order to experimentally dissect the transduction cascade. Furthermore, they would allow systematic pharmacological interference with the cAMP-dependent ORN subset, which is not possible so far.
1 Introduction

1.2 The effect of endocannabinoids on olfaction

Olfaction influences the behavior of animals as well as humans. If an olfactory cue is caught a distinct behavioral pattern might be elicited. But also vice versa, the current physiological state has an impact on olfactory processing. E.g. it is everyday experience that olfactory stimuli that are attractive before food intake may become neutral or even aversive afterwards. Even at the most peripheral state of the olfactory system, in the OE, hormones and transmitters modulate processing of odors. Thus, a single ORN cannot be considered as a static unit. Equal inputs may result in varying output, depending on the physiological state of the organism, and thus, the tuning of ORNs.

In the following section I will present a novel mechanism, by which odor responses are affected in a very fascinating way: the cannabinoid modulation. Then I will introduce some endogenous modulatory systems, which are known to exist in the OE and describe their function. In the last two sections I will give some physiological and biochemical facts about the endocannabinoid system.

1.2.1 Modulation of olfactory processing in the OE

Cannabinoids

The endocannabinoid system is an endogenous signaling system and affects multiple metabolic functions. The name is derived from the cannabis plant *Cannabis sativa* because of the cannabimimetic actions of its major active compound Δ9-tetrahydrocannabinol, which belongs to the group of cannabinoids (Gaoni and Mechoulam, 1964; Adams and Martin, 1996). The endocannabinoid system comprises cannabinoid (CB) receptors, their endogenous ligands, called endocannabinoids, and the proteins involved in the synthesis and degradation of these ligands (Mackie, 2008; Howlett *et al.*, 2002).

Recently, cannabinoid modulation of olfactory processing was observed. First, CB1 receptor mRNA was detected in the olfactory placodes of *Xenopus laevis* tadpoles (Migliarini *et al.*, 2006). One year later CB1 receptors could be localized to
the proximal dendrites of ORNs (Czesnik et al., 2007). They showed that CB1 receptor-specific antagonists modulate odor-evoked Ca$^{2+}$ changes in ORNs. Responses to odors were reduced and delayed. The delay was up to several seconds, which is an exceptional phenomenon and was not observed for other modulatory systems before. Since the CB1 receptor is expressed in ORNs and since ORNs can be modulated by cannabinoids, the existence of the endocannabinoid system in the OE seems very probable. This raises questions, which were not addressed as yet: Are endocannabinoids produced in the OE and if yes, which is the physiological trigger for endocannabinoid release? Is the odor sensitivity affected by endocannabinoids? And finally, how do endocannabinoids act?

**Other modulatory substances**

The modulation of olfactory processing is influenced by hormones involved in energy metabolism, like orexin and leptin. Orexin, an orexigenic peptide, is produced in the hypothalamus and acts on feeding and sleep. Leptin is secreted peripherally by adipocytes and acts in an anorexigenic manner. Additionally, both substances are synthesized locally in the OE (Caillol et al., 2003; Baly et al., 2007). Orexin may modulate transduction via binding to its receptors on ORNs and sustentacular cells (Caillol et al., 2003) and leptin decreases odor-induced transduction currents and receptor potentials upon binding to leptin receptors (Savigner et al., 2009), which are located on a subpopulation of ORNs (Baly et al., 2007). Cerebroventricular injection of orexin results in an increased olfactory sensitivity whereas leptin injection decreases sensitivity (Julliard et al., 2007).

ATP as well as dopamine act in a neuroprotective way in the olfactory system. ATP is thought to be released following noxious stimuli in the OE and reduces odor responsiveness. Vice versa, purinergic receptor antagonists have been shown to increase odor-evoked [Ca$^{2+}$]$_i$ transients (Hegg et al., 2003). Dopamine was also postulated to act in a neuroprotective manner in the OE (Hegg and Lucero, 2004; Féron, 1999). Stimulation of dopamine receptors on ORNs modulates hyperpolarization activated currents (Vargas and Lucero, 2002) and voltage-gated Ca$^{2+}$ channels (Hegg and Lucero, 2004).

Besides the mechanisms described so far, the gonadotropin releasing hormone modulates odor sensitivity in a season-dependent manner (Eisthen et al.,
2000). In addition, the neuroregenerative substances insulin like growth factor (Suzuki and Takeda, 2002; Mathonnet et al., 2001) and neuropeptide Y (Montani et al., 2006) act in the OE.

### 1.2.2 Physiological role of the endocannabinoid system

Appetite stimulation is probably the best-known effect of cannabis use. The ability of the endocannabinoid system to control appetite, food intake and energy balance has recently been described (Matias and Di Marzo, 2007; Osei-Hyiaman et al., 2006; Horvath, 2006). Additionally, selective inverse agonists of CB1 receptors reduce weight and can be used for the treatment of obesity (Kirkham and Tucci, 2006; Engeli, 2008). At the central nervous system level it has been well described that the endocannabinoid system plays a dual role in the regulation of food intake as well as in the homeostatic and non-homeostatic (or hedonic) energy regulation (Matias et al., 2008). Furthermore, it was shown that metabolic functions are controlled by endocannabinoids by acting on peripheral tissues, such as adipocytes, hepatocytes, and the gastrointestinal tract (Pagotto et al., 2006).

Another effect of cannabis consumption is an increased risk of developing a psychosis (Semple et al., 2005). This indicates that a disregulated endocannabinoid system may promote the development of e.g. depression, anxiety or schizophrenia. In fact, the CB1 receptor level in the prefrontal cortex of depressed suicide victims is elevated (Hungund et al., 2004), cannabinoid agonists have an anxiolytic action (Viveros et al., 2005), and schizophrenic patients have an increased CB1 receptor density in various brain regions as well as an increased endocannabinoid level in the cerebrospinal fluid (Zavitsanou et al., 2004; Giuffrida et al., 2004).

Furthermore, endocannabinoids also influence the perception of pain (Richardson et al., 1998), addiction (Scherma et al., 2008), stress (Gorzalka et al., 2008), neuroprotection (Galve-Roperh et al., 2008), and a variety of other functions.
1.2 The effect of endocannabinoids on olfaction

1.2.3 Biochemical aspects of the endocannabinoid system

Cannabinoid receptors

There are two major receptors which belong to the endocannabinoid system: CB1 and CB2 receptors (Matsuda et al., 1990; Munro et al., 1993). In addition to that, the orphan G protein-coupled receptor GPR55 was recently described to be targeted by a number of cannabinoids (Lauckner et al., 2008). The most abundant CB receptor in the central nervous system is CB1. This suggests that this receptor is responsible for the psychoactive effect of cannabinoids and the physiological actions of endocannabinoids (Elphick and Egertová, 2001).

The CB1 receptor belongs to the family of G protein-coupled receptors (Matsuda et al., 1990) and its distribution was first mapped by using the radiolabeled synthetic cannabinoid [3H]CP55,940. Using current techniques CB1 receptors could be localized in high quantity in the olfactory bulb and the hippocampus, in several parts of the striatum and its target nuclei, and the cerebellar molecular layer. Moderate CB1 receptor levels were identified (Figure 3) in other forebrain regions and in a few nuclei in the brain stem and the spinal cord (Herkenham et al., 1990; Herkenham et al., 1991). The CB1 receptor was also found in peripheral tissues like adipose tissue, liver, skeletal muscle, gastrointestinal tract, pancreas, thyroid gland, and adrenal gland (Pagotto et al., 2006; Demuth and Mollemann, 2006; Juan-Picó et al., 2006).

Endocannabinoids and their metabolism

The first endocannabinoid which was identified is N-arachidonoyl ethanolamide (or anandamide; Devane et al., 1992). It is a partial agonist for both CB receptors (Sugiura et al., 2002). In contrast, 2-arachidonoylglycerol (2-AG), which is found in much higher concentrations than anandamide in the brain (Sugiura et al., 2006), acts as a full agonist at the CB1 and CB2 receptors (Sugiura et al., 2002). Both endocannabinoids are produced on demand. This can be triggered by activation of Gq/11 protein-coupled receptors, by rising [Ca^{2+}] to the millimolar range, or by activation of Gq/11 protein-coupled receptors together with a moderate [Ca^{2+}] increase (Hashimotodani et al., 2007; Kano et al., 2009). The synthesis pathways (Figure 4) for anandamide and 2-AG are mediated by N-acylphosphatidylethanolamide-
Figure 3: Distribution of CB1 receptors in the central nervous system. The overall distribution of CB1 receptors in (A) sagittal and (B, C) frontal brain sections of mice was visualized by an immunolabeling with an antibody against the mouse CB1 receptor (Fukudome et al., 2004). CB1 immunoreactivity is highest along striatal output pathways, including the substantia nigra pars reticulata (SNR), globus pallidus (GP), and entopeduncular nucleus (EP). High levels are also observed in the hippocampus (Hi), dentate gyrus (DG), and cerebral cortex, such as the primary somatosensory cortex (S1), primary motor cortex (M1), primary visual cortex (V1), cingulate cortex (Cg), entorhinal cortex (Ent), basolateral amygdaloid nucleus (BLA), anterior olfactory nucleus (AON), caudate putamen (CPu), ventromedial hypothalamus (VMH), and cerebellar cortex (Cb). Scale bars 1 mm. Figure and figure subtitle modified from Kano et al., 2009.
1.2 The effect of endocannabinoids on olfaction

Anandamide and 2-AG are arachidonic acid derivates. Anandamide is synthesized by N-acyltransferase (NAT) and NAPE-PLD and degraded by FAAH, 2-AG is produced by phospholipase C (PLC) and DAGL and degraded by MAGL. The endocannabinoid membrane transporter (EMT) facilitates endocannabinoid release and uptake. The chemical structures of anandamide and 2-AG are indicated under the scheme. Figure from Di Marzo et al., 2004, see there for further information.
1 Introduction

hydrolyzing phospholipase D (NAPE-PLD) and diacylglycerol lipase (DAGL), respectively. Both endocannabinoids act extracellularly at CB receptors. For degradation, they are transported into cells, where they are primarily catabolized by the enzymes fatty acid amide hydrolase (FAAH) and monoacylglycerol lipase (MAGL), respectively (Basavarajappa, 2007; Kano et al., 2009).

Endocannabinoid signaling

Eight years ago, endocannabinoids have been found to act as retrograde messengers. They are released by postsynaptic neurons and modulate the release of inhibitory and excitatory neurotransmitters upon binding to CB1 receptors localized on presynaptic membranes. Besides this well established mechanism of depolarization-induced suppression of inhibition or excitation, endocannabinoids mediate synaptic plasticity and excitability (Kano et al., 2009; Howlett et al., 2004).

The CB1 receptor signaling is very diverse and seems to depend on the identity of the stimulating agonist as well as on the target tissue and the cell type (Howlett et al., 2002; Di Marzo et al., 2004; Demuth and Mollemann; 2006). In the following I will give an overview about the most abundant and best described signaling pathways (Figure 5).

Upon CB1 receptor stimulation, signal transduction pathways involving G<sub>i0</sub> or G<sub>S</sub> proteins can be activated (Howlett et al., 2005). Rhee et al. (1998) demonstrated that adenylate cyclase isoforms 1, 3, 5, and 8 were inhibited by CB1 receptor activation, and that the subtypes 2, 4, and 7 produced cAMP upon receptor stimulation. The G proteins coupled to CB1 receptors mediate a multitude of effects. One important aspect is the modulation of ion channels. Thus, activation of the CB1 receptor may result in the activation of K<sub>ir</sub> and A-type K<sup>+</sup> channels (McAllister et al., 1999; Childers and Deadwyler, 1996), and in the inhibition of voltage-gated L- (Gebremedhin et al., 1999; Straiker et al., 1999), N- (Mackie and Hille, 1992; Huang et al., 2001) and P/Q-type Ca<sup>2+</sup> channels (Mackie et al., 1995; Hampson et al., 1998). Furthermore, several intracellular kinases, like the protein kinase A and the mitogen-activated protein (MAP) kinase (Bouaboula et al., 1995; Galve-Roperh et al., 2002), are of particular importance for CB1 signaling, because these proteins alter gene expression, which then affects multiple cellular functions.
1.2 The effect of endocannabinoids on olfaction

What makes signaling of CB1 receptors even more complex is that these receptors form homo- and heterodimers (Mackie, 2005) and that they cross-talk with various other signaling systems, e.g. D2 receptors (Marcellino et al., 2008), opiate receptors (Robledo et al., 2008), and type1 orexin receptors (Hilairet et al., 2003).

**Figure 5: CB1 receptor signaling.** The activation of CB1 receptors leads to the stimulation of G proteins that in turn modulate various ion channels. In addition, several intracellular kinases are stimulated, which then modulate gene expression. Note that the figure shows only some of the known intracellular signaling events. Figure from Di Marzo et al., 2004. Abbreviations: AC: adenylate cyclase, PKA: protein kinase A.
1.3 Goal of the thesis

Olfactory disorders are associated with a variety of diseases that can hardly be medicated. In order to establish new drug therapies, it is substantially important to understand the underlying principles of olfaction. This thesis is subdivided into two parts, covering topics of basic research of olfaction:

1. At least two subsets of olfactory receptor neurons exist in the olfactory epithelium regarding odorant transduction mechanisms. One subset transduces odors using the cAMP-dependent transduction cascade, whereas the other subset uses a cAMP-independent transduction mechanism. In order to gain knowledge about, e.g., odor coding mechanisms, it would obviously be important, to be able to interfere with one of these subsets at the level of the generator current. This would make it possible to experimentally dissect the transduction of odors. Recently, the styryl dye FM1-43 was shown to stain olfactory receptor neurons, and to inhibit several cation channels of sensory cells. Therefore, I speculated that FM1-43 might be a promising candidate for blocking olfactory generator channels and examined this hypothesis. In this thesis, it will be shown that only a subset of the olfactory receptor neurons can be labeled with FM1-43. The identity of this subset will be characterized in detail. Furthermore, the effect of FM1-43 on the generator channel of this subset and on odor-induced responses will be examined.

2. The search for food as well as the subsequent food intake is known to be guided by the sense of smell, and it has been suggested that the feeding state modulates the olfactory sensitivity. However, the underlying mechanisms responsible for the functional interaction between olfaction and food intake are as yet poorly understood. It is well-documented that the endocannabinoid system is important for energy homeostasis and nutrition at central stages. The endocannabinoid system may therefore functionally link the feeding state and the olfactory sensitivity of an animal. Indeed, it was recently shown that cannabinoids act on olfactory receptor neurons. In this work, I will detect the endocannabinoid that acts in the olfactory epithelium and the cell types that
produce this substance. Moreover, the trigger for endocannabinoid release in
the olfactory epithelium, the effect on odorant detection thresholds as well as
the effector of the cannabinoid receptor will be investigated. Finally, the
existence of receptors for other modulatory substances besides endocan-
nabinoids in the olfactory epithelium will be examined.
2 Materials and Methods

2.1 *Xenopus laevis* tadpoles as the experimental model

*Xenopus laevis*, the South African clawed frog, belongs to the order “Anura” of the amphibians. The natural occurrence of this species is limited to Africa south of the Sahara. Because of its tremendous use as an animal model for many scientific purposes *Xenopus laevis* now occupies areas all over the world. This frog lives in the mud at the bottom of warm and stagnant water and ponds. It is predominantly crepuscular and nocturnal. The adult *Xenopus laevis* is counted among the scavengers, whereas the larvae feed mainly on algae. To locate food this species relies mostly on its sense of smell (Avila and Frye, 1978; Nieuwkoop and Faber, 1994).

The olfactory placode of larval animals becomes distinct at stage 23 as a thickening of the sensorial layer of the ectoderm (Klein and Graziadei, 1983). The olfactory organ begins to segregate into the principal cavity and the vomeronasal organ at stage 37/38 (Nieuwkoop and Faber, 1994). Synapses of ORN axons in the olfactory bulb also appear at this stage (Byrd and Burd, 1991). Mature ORNs could be specifically stained and olfactory receptor mRNA could be detected at stage 45 (Hansen *et al.*, 1998). A third sensory chamber, the middle cavity, forms and expands during metamorphosis. At the same time, the principal cavity is remodeled into the principal cavity of the adult animal (Hansen *et al.*, 1998; Nieuwkoop and Faber, 1994).

Since *Xenopus laevis* is totally aquatic as larva, the principal cavity of the tadpole is exposed to water-borne odorants, and after metamorphosis to airborne odorants. The middle cavity and the vomeronasal organ are always exposed to waterborne odorants (Freitag *et al.*, 1998). The vomeronasal organ detects pheromones (Halpern, 1987). The OE of *Xenopus laevis* consists of three cell types (Figure 6): ORNs (red), sustentacular cells (green, blue nuclei), and basal cells (blue nuclei shown at the basal side of the OE). Larval animals have ORNs bearing either cilia or microvilli, and sustentacular cells which are either ciliated or secretory. After
metamorphosis, the principal cavity comprises only ciliated ORNs and secretory sustentacular cells (Hansen et al. 1998).

In this thesis, *Xenopus laevis* tadpoles were used as the experimental model, because this aquatic animal is an excellent tool to study the olfactory system. First of all, it is easy to house adult frogs in the laboratory and every six to eight weeks spawning can be induced (Elinson, 2001). *Xenopus laevis* is a poikilothermal animal, and all experiments can be performed at room temperature. Slice preparations of the olfactory system of the tadpole are easy to produce, because the animals are essentially transparent with only a few melanocytes, and they have no cribriform plate between the OE and the olfactory bulb.

**Figure 6: Immunohistochemical staining of the OE of Xenopus laevis larvae.** Olfactory receptor neurons in a slice of the OE were stained with a biocytin-streptavidin backtrace (red). In green a cytokeratin-like-immunoreactivity of the sustentacular cells is shown. Sustentacular cells form a tightly packed layer on the apical side of the OE and their processes extend across the OE that terminate in endfeet-like structures at the basal level of the OE. All cell nuclei are stained with DAPI (blue). Scale bar: 15 µM. Figure kindly provided by T. Hassenklöver.
Adult frogs were purchased from Kaehler (Hamburg, Germany) and Nasco (USA) and held in aquaria with a water temperature of 20 °C. They were fed with Pondstick food (Tetra Pond, Melle, Germany). For inducing breeding, frogs were separated by gender and repeatedly injected with human chorionic gonadotropin (Sigma, Deisenhofen, Germany) subcutaneously. Then, breeding pairs were housed together overnight, and on the following day the embryos were obtained and kept in separate aquaria (water temperature 20 °C). The tadpoles were fed with algae (Dose Aquaristik, Bonn, Germany). For all experiments in this thesis, tadpoles of developmental stages 51 to 54 (Figure 7; Nieuwkoop and Faber, 1994) were used.

Figure 7: *Xenopus laevis* tadpole. (A) Larval *Xenopus laevis* are shown. (B) The olfactory system is marked by a black rectangle. The olfactory mucosa is located most anterior of the head. The ORN axons of the olfactory nerves terminate in the glomeruli in the olfactory bulb, the anterior part of the brain. All tadpoles are from stage 54. Figures kindly provided by T. Hassenklöver and I. Manzini.
2.2 *In vivo* labeling of ORNs with FM1-43

To stain ORNs with FM1-43 (N-(3-triethylammoniumpropyl)-4-(4-(dibutylamino)-styryl)pyridinium dibromide, Figure 8), living tadpoles were transferred into distilled water for 5 min and then incubated for 7 min (standard staining) or 1 min and 15 s (light staining) in 10 ml 2 µM FM1-43 (Molecular Probes, Leiden, the Netherlands) dissolved in distilled water. Afterwards the tadpoles swam again briefly in distilled water. In some experiments, where the impact of certain substances on the staining efficiency was investigated, 2 mM CaCl₂, 1 mM MgCl₂, 200 µM LY-83,583 or 1 mM amiloride was added to the solution that contained FM1-43. In these cases the exposure time in the respective incubation solution was 7 min.

Subsequently the animals were anaesthetized for OE slice preparation and viewed by using a laser-scanning confocal microscope attached to an inverted microscope (LSM 510) with 10x or 40x objectives. The confocal pinhole was set to 120-150 µm to exclude fluorescence detection from more than one cell layer. Fluorescence images (excitation at 488 nm; emission > 505 nm) of the OE were acquired together with a pseudo bright-field image for orientation in the tissue.

FM1-43 stained tadpoles were also used for OE slice preparation with subsequent double labeling for [Ca²⁺]ᵢ imaging experiments.

2.3 Tissue preparations

2.3.1 OE slice preparation

Slices were prepared of animals, which were prestained with FM1-43 or non-stained. For this purpose, the tadpoles of *Xenopus laevis* were chilled in a mixture of ice and water and decapitated, as approved by the University of Göttingen Committee for Ethics in Animal Experimentation. A block of tissue containing the OE, the olfactory nerves, and the brain was cut out and kept in bath solution (Table 2). The tissue was then glued onto the stage of a vibratome (VT 1200S; Leica, Bensheim, Germany) and cut horizontally into 130 to 150 µm thick slices.
For imaging soma $[\text{Ca}^{2+}]_i$, tissue slices were incubated with 200 µl of bath solution containing 50 µM fluo-4 AM (Molecular Probes, Leiden, The Netherlands) and 50 µM MK571 (Alexis Biochemicals, Grünberg, Germany). Fluo-4 AM was dissolved in DMSO (Sigma, Deisenhofen, Germany) and Pluronic F-127 (Molecular Probes). The final concentrations of DMSO and Pluronic F-127 did not exceed 0.5 % and 0.1 %, respectively. To avoid multidrug resistance transporter mediated destaining of the slices, MK571, a specific inhibitor of the multidrug resistance-associated proteins, was added to the incubation solution (Manzini et al., 2003b). After incubation at room temperature for 30 min, the tissue slices were put under a grid in a recording chamber and placed on the microscope stage of an Axiovert 100M (Zeiss, Jena, Germany) to which a laser scanning unit (LSM 510; Zeiss) was attached. Before starting the $\text{Ca}^{2+}$ imaging experiments, the slices were rinsed with bath solution for at least 5 min.

2.3.2 Olfactory bulb whole mount preparation

For imaging $[\text{Ca}^{2+}]_i$ of glomeruli, ORNs were traced via fluo-4 dextran 10 kDa by electroporation in the OE. For this purpose, larval *Xenopus laevis* were anesthetized in 0.02 % MS-222 (Sigma). Crystals of fluo-4 dextran, potassium salt, 10 kDa
(Invitrogen) were inserted into the nasal cavities, where it dissolved in the residual water. Subsequently, two platinum electrodes of 250 µm diameter and 3 mm interspace interval were placed into the nasal cavities and the dye was electro-porated by application of 30 V twelve times with alternating polarity. The animals were kept in a water tank for one to three days until experiments were performed.

Tadpoles were chilled in a mixture of ice and water and decapitated, as approved by the University of Göttingen Committee for Ethics in Animal Experimentation. A block of tissue containing the OE, the olfactory nerves, and the brain was cut out and kept in bath solution (Table 2). The tissue surrounding the ventral part of the olfactory bulb was removed and the whole-mount preparation was put under a grid in a recording chamber and placed on the microscope stage of an Axiovert 100M (Zeiss, Jena, Germany) to which a laser scanning unit (LSM 510; Zeiss) was attached. The tissue was rinsed with bath solution for at least 5 min before the experiment was started.

2.4 Conventional and advanced \([\text{Ca}^{2+}]_i\) imaging and patch clamp recordings

2.4.1 \([\text{Ca}^{2+}]_i\) imaging of odor-induced responses of ORNs and glomeruli with confocal microscopy

For \([\text{Ca}^{2+}]_i\) imaging of odor-induced responses of ORNs, fluo-4 stained tissue slices were produced. Glomeruli were imaged using whole mount preparations prestained with fluo-4 dextran.

\([\text{Ca}^{2+}]_i\) was monitored using a laser-scanning confocal microscope (LSM 510, Zeiss). The confocal pinhole was set to approximately 120 µm / 300 µm to exclude fluorescence detection from more than one cell layer / glomerulus. Fluorescence images (excitation at 488 nm; emission > 505 nm for fluo-4 stained OE slices and glomeruli; emission from 505 to 530 nm and > 560 nm for fluo-4 and FM1-43 doublestained OE slices, respectively) of the OE were acquired in the range of 1.02 to 2.03 Hz, with three to ten images taken as control images before the onset of odor
delivery. The fluorescence changes $\Delta F/F$ of fluo-4 were calculated for individual ORNs / glomeruli as $\Delta F/F = (F_1 - F_2) / F_2$, where $F_1$ was the fluorescence averaged over the pixels of an ORN soma / glomerulus, while $F_2$ was the average fluorescence of the same pixels prior to stimulus application, averaged over five images. A response was assumed if the following two criteria were met: (i) the first two intensity values after stimulus arrival at the mucosa, $\Delta F/F(t_1)$ and $\Delta F/F(t_2)$, had to be larger than the maximum of the prestimulus intensities; (ii) $\Delta F/F(t_2) > \Delta F/F(t_1)$ with $t_2 > t_1$.

Data analysis was performed with Matlab (Mathworks, USA).

### 2.4.2 Uncaging of cAMP in ORNs viewed with confocal microscopy

In order to observe the effect of cAMP uncaging on $[\text{Ca}^{2+}]_i$ in FM1-43-loaded ORNs, tadpoles were stained with the dye as described and OE slices were prepared. Afterwards the tissue was incubated in 200 µl of $\text{Ca}^{2+}$ indicator rhod-2 AM solution (50 µM rhod-2 AM (Molecular Probes, Leiden, The Netherlands) dissolved in DMSO (0.5 %) and Pluronic F-127 (0.1 %) and 50 µM MK571) at room temperature for 30 min. The tissue slices were placed under a grid in a recording chamber and positioned on the microscope stage of an Axiovert 100M to which a laser scanning unit was attached. A glass fiber (HCG-M0200T 200 µm, Laser Components) coupled to a 378 nm diode laser (iPulse, Toptica Photonics) was positioned next to the OE. The slices were incubated with 100 µM DMNB-caged cAMP (Invitrogen, stock solution: 20mM in DMSO) for 15 min. Stimulation of ORNs was performed by a 10 ms laser pulse (378 nm, 16 mW).

After FM1-43-loaded ORNs were identified, rhod-2 fluorescence was monitored using a pinhole diameter of approximately 120 µm. Images (excitation at 543 nm; emission > 560 nm) of the OE were acquired in the range of 1.02 to 2.03 Hz, with three to ten images taken as control images before the onset of odor delivery. The fluorescence changes $\Delta F/F$ were calculated for individual ORNs. Data analysis was performed with Matlab.
2.4.3 \([\text{Ca}^{2+}]_i\) dendrite imaging in tissue slices with a fast scanning line illumination microscope

For fast imaging of dendrite \([\text{Ca}^{2+}]_i\), slices were placed under a grid in a recording chamber and viewed by a 63x water immersion objective mounted to a custom-built line illumination microscope described by Junek et al. (2009). Patch pipettes were filled with 4 µl fluo-4 containing pipette solution (Table 3). After establishing the whole-cell mode fluo-4 diffused into the cell. Subsequently, stacks of images of the stained ORN were obtained every 30 s. Each image stack comprised 20 images and was acquired within 328 ms.

2.4.4 Patch-clamp recordings of the CNG current

For patch-clamping the slices were placed under a grid in a recording chamber and viewed by using Nomarski optics (Axioskop 2; Zeiss, Göttingen, Germany). Patch pipettes with a tip resistance of 6-10 MΩ were pulled from borosilicate glass with a 1.8 mm outer diameter (Hilgenreiner, Malsfeld, Germany) by a two-stage pipette puller (PC-10, Narishige, Japan) and filled with 4 µl cAMP and cGMP containing pipette solution (Table 3). Pulse protocols for data acquisition were written in C. Voltage pulses were delivered from a microcontroller (Schild et al., 1996) to a D/A converter and then to the patch-clamp amplifier (EPC7; List, Darmstadt, Germany) in order to assess the impedance in the on-cell and whole-cell configurations. The data were digitized online.

After establishing the on-cell configuration in bath solution the holding potential was set to 0 mV. The responsiveness of the patch-clamped cell was tested by stimulating the ORN with 50 µM forskolin dissolved in bath solution. Subsequently, the holding potential was set to -70 mV and the external solution was replaced by \(\text{Ca}^{2+}\) and \(\text{Mg}^{2+}\) free bath solution (Table 2) with or without 10 µM FM1-43. Directly after establishing the whole-cell configuration by shortly applying negative pressure to the patch pipette an inward current was induced. Evaluation of the current traces was performed in Matlab.
2 Materials and Methods

2.4.5 Bathing conditions and stimulus application

For all experiments the recording chamber was perfused by gravity feed from a syringe through a funnel applicator (Schild, 1985; Manzini et al., 2002) with various bath solutions (Table 2). The tip of the applicator was placed in front of the OE. The change of the external solution was performed by stopping the influx of one bath solution and starting the influx of another bath solution into the funnel applicator.

A mixture of amino acids (Manzini et al., 2002; Caprio and Byrd, 1984; Iida and Kashiwayanagi, 1999), amines (Carr and Derby, 1986; Carr et al., 1990; Rolen et al., 2003; Gliem et al., 2009), bile acids (Kang and Caprio, 1995; Sato and Suzuki, 2001) and alcohols (Altner et al., 1977; Tinsley and Kobel, 1996) are known to be odorants for aquatic species and were used as odors. A mixture of 19 amino acids or single amino acids (arginine, lysine and methionine), as well as mixes of amines, bile acids, and alcohols were used as odors. The odors were dissolved in bath solution (stocks of 10 mM or 25 mM, see Table 4) and used at a final concentration of 0.2 µM to 100 µM in all of the experiments. Stimulus solutions were prepared immediately before use and were pipetted directly into the funnel for bath perfusion without stopping the flow. Outflow was through a syringe needle placed close to the OE. The time course of stimulus arrival at the OE was simulated by applying the fluorescent dye avidin AlexaFluor-488 as a dummy stimulus and by measuring the fluorescence after avidin AlexaFluor-488 application to the funnel. The delay of stimulus arrival caused by the syringe, i.e., from pipetting into the funnel to concentration increase in the OE, was approximately 2 s. The minimum interstimulus interval between odorant applications was 2 min.

2.5 Molecular biology experiments

2.5.1 Single-cell reverse transcription (RT) PCR

Tissue slices were visualized using a 40x water immersion objective mounted to an Axioscop 2 microscope. Patch pipettes were filled with 4 µl pipette solution (Table 3).
2.5 Molecular biology experiments

Cells were identified as ORNs and sustentacular cells based on their morphology. After the formation of a gigaseal, negative pressure was applied to the pipette and the whole cell configuration was established (Hamill et al., 1981). Olfactory receptor neurons showed spontaneous spiking activity in the on-cell mode and typical voltage-gated Na$^+$ and K$^+$ currents in the whole cell configuration. Sustentacular cells typically show no electric activity. Cell cytoplasm was harvested under visual and resistance control by applying gentle suction to the patch pipette.

Cells fulfilling these physiological criteria and whose seals remained intact during harvesting were used for reverse transcription with a modified protocol of the SuperScript$^\text{TM}$ III First-Strand Synthesis System for RT PCR (Invitrogen). The content of the pipette was immediately expelled into a tube containing 5 ng random hexamers, 40 U RNasin Plus RNase Inhibitor (Promega), 1 mM dNTP mix, and DEPC water. The mixture was heated to 65 °C for 5 min and cooled on ice for at least 1 min. Next, reverse transcription was performed by adding 1x RT buffer, 5 mM MgCl$_2$, 10 mM DTT, 2 U RNaseOUT, and 10 U SuperScript III RT and incubating in a thermocycler (T-Personal, Biometra) at 25 °C (10 min), 50 °C (50 min), 85 °C (5 min), and chilled on ice. RNA was degraded by adding 1 µl RNase H and incubating for 20 min at 37 °C. Negative control reactions without SuperScript III RT were also performed.

The cDNA produced in one single cell RT was split in four tubes and served as the template for PCR. The reactions were performed according to the manual of the FastStart Taq DNA Polymerase (Roche). In brief, the reaction mix contained 200 nM specific forward and reverse primers for OMP1 (Rössler et al., 1998), CYTII, DAGL$\alpha$, or DAGL$\beta$ (primer sequences in Table 1), 200 µM dNTPs, 1x PCR buffer, and 2 U FastStartTaq DNA Polymerase. The reaction was activated at 95 °C for 5 min and underwent 40 cycles of a temperature protocol of 30 s at 95 °C, 30 s at 58 °C, and 45 °C at 72 °C. After a final extension of 7 min at 72 °C the PCR products were run on a 2 % (w/v) agarose gel in tris acetate EDTA (TAE) buffer containing ethidium bromide (Sigma) and visualized under UV-light (UVsolo, Biometra).
2.5.2 Real-time PCR

Tadpoles were exposed to four different nutritious states (n = 7): In one condition animals were food-deprived for 6 h (group A$_{6h}$) or 12 h (group A$_{12h}$) in another animals were food-deprived for 6 h and overfeed for 2 h with 1 g shredded algae (Dohse Aquaristic, Millipore; group B$_{6h}$) per 1 l water or 12 h food-deprived for 12 h and overfeed for 2 h (group B$_{12h}$). As control condition animals were overfeed for 2 h (group C).

RNA isolation and cDNA synthesis

Olfactory epithelia of four animals per condition were cut out of the tissue and stored in liquid nitrogen until RNA isolation. Total RNA was isolated with the TRIzol method (Invitrogen) according to the manufacturer’s protocol and DNA-contaminations were removed by subsequent DNase I treatment (DNase I recombinant, RNase-free, Roche). The RNA quality and quantity was analyzed with the microfluidics-based electrophoresis system Agilent 2100 Bioanalyzer (Agilent Technologies). Reverse transcription was performed from 1 µg RNA with the iScript cDNA Synthesis Kit from BioRad as described in the manual.

cDNA Quantification

Quantification of DAGL$\alpha$ and $\beta$ RNA was performed using the iQ SYBR Green Supermix (BioRad) on an iQ5 real-time PCR detection system (BioRad) according to the manufacturer's instructions. The ATPase F$_0$F$_1$ (primer sequences see Table 1) was used as an internal control. The general PCR conditions were as follows: polymerase activation at 98 °C for 30 s followed by 40 cycles of denaturation at 94 °C for 1 s, annealing at 58 °C for 15 s, and extension at 72 °C for 1 s. After the amplification a melt curve analysis verified the formation of the single desired PCR products. The relative gene expression ratios (Kubista et al., 2006) were determined and normalized for control conditions. Confidence intervals were calculated by determining the standard deviation of the logarithmized ratios followed by exposing the left and right borders.
2.5.3 Conventional PCR

For the analysis of gene expression of *Xenopus laevis*, *Rattus norvegicus* and *Homo sapiens* mRNA was analyzed. The OEs of three rats were excised. For *Xenopus laevis* tadpoles, ten OEs were pooled for all three samples. mRNA isolation and reverse transcription were accomplished as described. Human cDNA was kindly provided by Thomas Hummel. For PCR, 200 ng of cDNA were mixed with PCR buffer, 200 µM nucleotide mix, 200 nM forward and reverse primers (Table 1), and 2 U FastStartTaq DNA Polymerase dissolved in PCR grade water as described in the manual (FastStart Taq DNA Polymerase, dNTPack, Roche, Mannheim, Germany). The samples were incubated in a thermocycler with the following PCR conditions: polymerase activation at 95 °C for 5 min followed by 30 cycles of denaturation at 95 °C for 30 s, annealing at 56 °C for 30 s, and extension at 72 °C for 45 s. After a final elongation at 72 °C for 7 min PCR products were run on a 2 % agarose gel in TAE buffer and ethidium bromide and visualized under UV light.
## 2 Materials and Methods

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<td>reverse primer</td>
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<td></td>
<td>product length [bp]</td>
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<tr>
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2.5 Molecular biology experiments

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<tr>
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**Homo sapiens**

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<th>Primer sequence (3' to 5')</th>
<th>Length (bp)</th>
</tr>
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<tr>
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**Table 1: Primer sequences.** All sequences listed are written from 5’ to 3’. The resulting length of the PCR products are indicated in base pairs (bp). All primers were purchased from Invitrogen. Abbreviations of gene names: OMP1: olfactory marker protein 1, CB1: CB1 receptor, D2A and B: dopamine receptor 2A and B, AdR1 and 2: adiponectin receptor 1 and 2, leptinR: leptin receptor, CB2: CB2 receptor, GPR55: G protein-coupled receptor 55, orexinR1 and 2: orexin receptor 1 and 2.
2.6 Solutions

2.6.1 External solutions

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</tr>
</tbody>
</table>

adjusted to pH 7.8 and 230 mOsmol/liter

Table 2: Composition of external solutions. The composition of all bath solutions used are listed in this table. All components were purchased from Sigma. Abbreviations: bs: bath solution, 0 Ca 0 Mg: Ca$^{2+}$ and Mg$^{2+}$ free bath solution, NMDG25/10/0: bath solution with substituted Na$^+$ (25/10/0 mM Na$^+$ and 73/88/98 mM NMDG, a Na$^+$ substitute).
2.6.2 Internal solutions

Stock solutions for Na$_2$-ATP (100 mM), Na$_2$-GTP (10 mM), cAMP (100 mM), and cGMP (10 mM) were prepared in HEPES solution (pH 7.8), fluo-4 potassium salt was dissolved in distilled water (10 mM).

<table>
<thead>
<tr>
<th></th>
<th>ps</th>
<th>cAMP/cGMP ps</th>
<th>fluo-4 ps</th>
</tr>
</thead>
<tbody>
<tr>
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<td>MgSO$_4$</td>
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<td>2 mM</td>
</tr>
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<td>K-gluconate</td>
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</tr>
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<td>0.2 mM</td>
<td>0.2 mM</td>
</tr>
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<td>1 mM</td>
<td>1 mM</td>
</tr>
<tr>
<td>Na$_2$-GTP</td>
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<td>0.1 mM</td>
</tr>
<tr>
<td>cAMP</td>
<td>-</td>
<td>1 mM</td>
<td>-</td>
</tr>
<tr>
<td>cGMP</td>
<td>-</td>
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<td>-</td>
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<td>fluo-4 K$^+$ salt</td>
<td>-</td>
<td>-</td>
<td>0.1 mM</td>
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</tbody>
</table>

adjusted to pH 7.8 and 190 mOsm/liter

**Table 3: Composition of internal solutions.** The composition of all pipette solutions are listed here. All components besides fluo-4 potassium salt (Invitrogen) were purchased from Sigma. Abbreviations: ps: pipette solution, cAMP/cGMP ps: cAMP and cGMP containing pipette solution, fluo-4 ps: fluo-4 potassium salt containing pipette solution.
### 2.6.3 Odors

<table>
<thead>
<tr>
<th>odor</th>
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<th>odor</th>
<th>concentration of stock solution</th>
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<td>amine mix</td>
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</tr>
<tr>
<td>L-alanine</td>
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<td>tyramine</td>
<td>25 mM</td>
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<tr>
<td>L-serine</td>
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</tr>
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<td>L-cysteine</td>
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<td>25 mM</td>
</tr>
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<td>10 mM</td>
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</tr>
<tr>
<td>L-glutamine</td>
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<td>25 mM</td>
</tr>
<tr>
<td>L-valine</td>
<td>10 mM</td>
<td>2-methylbutylamine</td>
<td>25 mM</td>
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<td>L-leucine</td>
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<td>25 mM</td>
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<td>L-lysine*</td>
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<td>alcohol mix</td>
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<td>L-histidine</td>
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<td>L-tryptophane</td>
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<td>terpineol</td>
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<td>bile acid mix</td>
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</tr>
<tr>
<td>taurocholic acid</td>
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**Table 4: Odors.** All odors were used as the indicated mixes or as the single amino acids arginine, lysine, and methionine (marked with *). All chemicals were purchased from Sigma and stock solutions were made in bath solution.
All odors were used at a final concentration of 100 µM dissolved in bath solution. Only the single amino acids arginine, lysine, and methionine were applied in various concentrations as indicated in the corresponding results section.

### 2.6.4 Drugs

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<tr>
<td>CE</td>
<td>Invitrogen</td>
<td>DMSO</td>
<td>10 mM</td>
<td>5 µM</td>
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</tbody>
</table>

Table 5: Drug solutions. The purchased drugs are listed with stock and working solutions. Abbreviations: CE: SE(5-(and-6)-carboxy eosin diacetate, succinimidy  ester).
3 Results

3.1 The styryl dye FM1-43 permeates and blocks CNG channels in olfactory receptor neurons of larval *Xenopus laevis*

3.1.1 FM1-43 stains a subset of ORNs

In a first set of experiments, living *Xenopus laevis* tadpoles were put into water containing the styryl dye FM1-43 (2 µM). Thereafter the animals were sacrificed and tissue slices were prepared from the OE. When the slices were viewed with a confocal laser scanning microscope, a large number of cells were stained in the entirety of their cytosol (Figure 9 A), whereas control slices showed no fluorescence (exposed to the same treatment but without FM1-43; Figure 9 B). For a better orientation the fluorescence images were overlayed with the corresponding transmission images scanned through wide-field optics. Figure 9 C shows the magnified rectangular area of B as a z-projection to illustrate the fine structure of the stained cells. Dendrites running to the surface of the OE, where cilia or microvilli issued from dendritic knobs, and axons running into the opposite direction to join the olfactory nerve unambiguously defined these cells as ORNs. Nuclei remained unstained. No staining at all was found in the vomeronasal organ (not shown).

3.1.2 FM1-43-stained ORNs rarely respond to odors

FM1-43 never stained the entire OE. It rather appeared to stain a certain subset of ORNs. To characterize the ORNs of this subset it was tried to test their sensitivity to amino acids, bile acids, amines, alcohols, and a mixture of all (100 µM for each substance). 156 out of 165 stained ORNs did not respond to any of the stimuli, which is in contrast to the high responsiveness of *Xenopus* tadpole ORNs as seen in previous studies (Manzini and Schild, 2004; Schild and Manzini, 2004). Only nine
ORNs were responsive to the mixture, one of them to alcohols and four to amines. Figure 10 A shows an FM1-43-loaded cell (red) in a fluo-4 stained slice (green) and Figure 10 B–H gives a typical example showing primarily two things. First, this ORN was sensitive to alcohols (Figure 10 D, G) but not to amino acids (Figure 10 C), bile acids (Figure 10 E) and amines (Figure 10 F) and second, the response amplitudes to both the stimulus mixture (Figure 10 B, H) and to alcohols (Figure 10 D, G) rapidly declined over time and then vanished. The facts that FM1-43 stained only a subset of ORNs and that most of the stained ORNs did not respond at all, while those few which initially did respond rapidly lost their responsiveness, suggested that the responsiveness of the stained ORNs was severely compromised by FM1-43.

**Figure 9:** FM1-43 is selectively internalized by a subset of ORNs. (A, B) The image shows OEs of tadpoles with and without FM1-43 incubation, respectively. (C) The z-projection and magnification of (A) illustrates the morphology of FM1-43-labeled ORNs by the cytosolic staining. Scale bars: (A, B) 50 µm, (C) 10 µm.
Figure 10: Odor-induced \([\text{Ca}^{2+}]\) transients of an FM1-43-stained ORN. (A) The image shows a doublestained OE slice with fluo4-AM (green) and a FM1-43-loaded ORN (red; image acquired at rest). (B-H) The \([\text{Ca}^{2+}]\) transients of the FM1-43-labeled ORN from (A) evoked by odorant mixtures (B, H), amino acids (C), alcohols (D, G), bile acids (E), and amines (F; 100 µM for each) are shown in chronological order. Scale bars: 10 s and \(\Delta F/F = 10\%\). The black lines indicate the application of the odorants.
3.1 FM1-43 permeates and blocks CNG channels

As FM1-43 uptake took place in the OE in vivo, it certainly occurred through the plasma membrane of the compartments exposed to the principal cavity, i.e., through cilia, microvilli, and/or dendritic knobs. Furthermore, as FM1-43 fluorescence was cytosolic and as it built up rapidly, FM1-43 permeated into the cytosol presumably via ion channels rather than via transport proteins. It was therefore checked whether CNG channels were permeable for FM1-43 using the well-known permeability properties of divalent ions in CNG channels as well as the effect of two non-specific blockers of CNG channels.

When CaCl₂ (2 mM; n = 5) or MgCl₂ (1 mM; n = 5) was added to the water during in vivo incubation with FM1-43, the fluorescence intensity of ORNs was reduced to almost zero (Figure 11 A (CaCl₂) and Figure 11 B (MgCl₂), control: Figure 11 C). This would be consistent with an uptake of FM1-43 through CNG channels as

---

**Figure 11: Block of FM1-43 labeling by cations.** Incubation of the tadpoles in FM1-43 solution with (A) 2 mM CaCl₂ or (B) 1 mM MgCl₂ almost completely blocked FM1-43 uptake. (C) Under control conditions many ORNs were labeled when living tadpoles were incubated in 2 µM FM1-43 solution. Scale bars: 200 µm.

---

3.1.3 FM1-43 is selectively internalized by CNG channels
Ca\(^{2+}\) has been reported to exert a permeation block in these channels (Frings et al., 1995).

If FM1-43 permeates through CNG channels its permeation should be affected by LY-83,583 or amiloride. When LY-83,583 (200 µM), which blocks CNG channels and the soluble guanylyl cyclase (Leinders-Zufall and Zufall, 1995), was added during dye incubation, the uptake of FM1-43 was completely blocked (Figure 12 A, n = 10; control: Figure 12 C). The presence of amiloride (1 mM), which blocks CNG channels, Na\(^{+}\) channels, T-type Ca\(^{2+}\) channels and several transporters (Benos, 1982; Zhuang et al., 1984; Tang et al., 1988; Frings et al., 1992), during incubation also reduced FM1-43 uptake dramatically (Figure 12 B, n = 8). These results suggest that CNG channels have a sizable permeability for FM1-43. The ORNs stained by FM1-43 may thus correspond to the subset of ORNs endowed with the canonical cAMP-transduction cascade.

The direct test of this hypothesis would be to evoke responses to cAMP in FM1-43-stained cells. Of course, this is conflicting with the hypothesis itself because FM1-43 would suppress the responses. It was tried to circumvent this problem by exposing the animals to FM1-43 for a relatively short time in order to have a

Figure 12: Block of FM1-43 labeling by unspecific CNG channel blockers. Incubation of the tadpoles in FM1-43 solution with the unspecific CNG channel blockers (A) LY-83,583 (200 µM) or (B) amiloride (1 mM) blocked FM1-43 uptake compared to control conditions (C). Scale bars: 200 µm.
3.1 FM1-43 permeates and blocks CNG channels

correspondingly weak staining and at least some CNG channels functional. In fact, under these conditions, the ORN staining with FM1-43 was rather faint, and when the cells were stimulated with forskolin they showed weak but clear and reproducible responses (Figure 13 A). Similar results were obtained in ten out of 13 cells (five slices). The three non-responding cells came all from the same slice. Uncaging of caged cAMP in FM1-43-loaded ORNs also resulted in a small, transient fluorescence increase of the Ca$^{2+}$ indicator dye rhod-2 (Figure 13 B; five out of five cells; three slices; performed together with E. Kludt).

Taken together, the block of FM1-43 uptake by divalent ions and by CNG channel blockers as well as the responses of faintly stained ORNs to forskolin and cAMP is consistent with the hypothesis that FM1-43 enters ORNs through CNG channels.

Figure 13: FM1-43-labeled ORNs are sensitive to forskolin and uncaging of cAMP. (A) Forskolin-evoked [Ca$^{2+}$]$_i$ transients and (B) [Ca$^{2+}$]$_i$ transients induced by uncaging of cAMP in individual FM1-43-stained ORNs are reproducible and have small amplitudes. Scale bars: (A) 20 s and $\Delta F/F = 10\%$, (B) 10 s and $\Delta F/F = 5\%$. The black line indicates the application of the forskolin and the black dot the time point of uncaging.
3.1.4 FM1-43 inhibits CNG currents

Patch-clamped ORNs in untreated OE tissue slices were first identified as cAMP-dependent or -independent by stimulation with forskolin in the on-cell mode of the patch clamp technique. Some ORNs responded to forskolin with a transient firing rate increase (Figure 14 A and B, upper traces), while others, presumably due to the lack of CNG channels, showed no response to forskolin (Figure 14 C, upper trace). In a second step of the experiment the same cells were recorded in the whole-cell mode, with cAMP and cGMP added to the pipette solution. The effect of the second messengers that diffuse from the pipette into the cell was observed either with (Figure 14 A) or without FM1-43 (Figure 14 B) added to the bath solution. Without any FM1-43 in the bath an inward current set in immediately after breakthrough (Figure 14 A, blue trace). To avoid, as much as possible, the activation of Ca\(^{2+}\)-activated Cl\(^{-}\) channels downstream the CNG channels, Ca\(^{2+}\) was omitted from the bath in these experiments, so that the recorded current was a current through CNG channels carried by Na\(^{+}\) ions. Its average amplitude was 213.8 +/- 21.2 pA (SEM; Figure 14 A, blue trace; n = 5). FM1-43 in the bath solution (10 µM) reduced the inward current in cAMP-dependent cells upon breakthrough to 54.5 +/- 31.6 pA (Figure 14 B, red trace; n = 6). In non-cAMP-dependent ORNs, cAMP and cGMP never had any effect on the current (Figure 14 C, blue trace; n = 4). An overview about the reduced CNG current amplitudes is given in Figure 14 D.

3.1.5 Extracellular FM1-43 in the OE reduces forskolin-induced responses of glomeruli

The previous experiment demonstrated that FM1-43 inhibits CNG channels, but the site of inhibition remains unclear. Therefore, the effect of extracellular FM1-43 on odor responses was investigated (experiment performed together with E. Kludt). Figure 15 A (black trace) shows a forskolin-application elicited [Ca\(^{2+}\)] transient in a glomerulus in the medial cluster of a typical bulb whole mount preparation. When FM1-43 (10 µM) was added to the bath solution, the amplitudes of the [Ca\(^{2+}\)] transients were reduced (red trace). This effect was reversible by washing FM1-43
3.1 FM1-43 permeates and blocks CNG channels

Figure 14: CNG currents are inhibited by FM1-43. (A, B, C) Cells were patch-clamped in the on-cell mode in bath solution. Forskolin induced an increased spike frequency in cAMP-dependent (A and B, black traces), but not in cAMP-independent ORNs (C, black trace). Subsequently, the bath solution was substituted with Ca²⁺- and Mg²⁺-free bath solution (0 Ca 0 Mg) and the whole cell mode was established with cAMP and cGMP in the pipette solution. This induced an inward current in cAMP-dependent ORNs (A, blue trace), no current was detected in cAMP-independent neurons (C, blue trace). (B, red trace) When FM1-43 was present in the Ca²⁺- and Mg²⁺-free bath solution, the amplitude of the inward current was reduced dramatically. (D) The current amplitudes of (A; n = 5) and (B; n = 6) are quantified in a bar graph. Scale bars: (A-C) 5 s and 50 mV or 50 pA. The black lines indicate the application of the forskolin.

out of the slice (grey trace; 11 glomeruli in 3 animals). Figure 15 B summarizes the obtained data. The mean amplitudes of the [Ca²⁺]i transients during FM1-43 wash-in and during wash-out are depicted normalized to those of the control experiment. The
amplitude was reduced to 0.58 ± 0.16 upon 10 µM FM1-43 in the bath compared to control conditions. Wash-out of the dye increased the amplitude to 0.67 ± 0.22 (Figure 15 B).

**Figure 15: Extracellular FM1-43 inhibits glomerular responses.** (A) Forskolin-evoked 
$[Ca^{2+}]_i$ transients of glomeruli in the medial cluster of a typical olfactory bulb whole mount preparation (black trace) were reduced upon FM1-43 in the bath (10 µM; red trace). Wash-out of the dye recovered the amplitude (grey trace; n = 11). (B) The amplitudes are quantified in the bar graph. Scale bar: (A) 10 s and $\Delta F/F = 20 \%$. The black line indicates the application of forskolin.
3.2 Endocannabinoid modulation in the olfactory epithelium

3.2 Modulation of processing in olfactory receptor neurons by the endogenous cannabinoid system

3.2.1 Localization of the endocannabinoid system in OE

Czesnik et al. (2007) proved that cannabinoids affect olfactory processing. This finding makes the presence of the endogenous cannabinoid system in the OE probable. In order to locate the endocannabinoid system in the OE, the mRNA content of the whole OE was analyzed for components of the endocannabinoid system with PCR. cDNA for the CB1 receptor, the 2-AG-catabolizing enzymes DAGLα and β and MAGL, and the anandamide-catabolizing enzymes NAPE-PLD and FAAH were detected (Figure 16). Thus, 2-AG and anandamide can be produced and act in the OE.

3.2.2 Suppression of 2-AG production reduces and delays odor-induced responses of ORNs

Endocannabinoids play a physiological role in the OE. When the CB1 receptors of ORNs are blocked, responses to odorants are diminished and delayed (Czesnik et al., 2007). This effect could be explained by assuming a tonic synthesis and action of endocannabinoids in the OE. This assumption was checked by blocking 2-AG synthesis using the DAGL blockers RHC80267 or orlistat. The superfusion with these drugs had two effects. They prolonged the delay and reduced the amplitude of responses of individual ORNs to odorants. The black traces in Figure 17 A and B show typical $[Ca^{2+}]_i$ responses upon application of a mixture of amino acids (100 µM) in two different ORNs taken from two different OE slice preparations. Superfusion of the slices with orlistat (50 µM, for 10 min) or RHC80267 (50 µM, for 12 min) diminished and delayed the $[Ca^{2+}]_i$ responses (Figure 17 A, B, respectively, red traces). This effect was highly reproducible (observed in 49 out of 49 cells, eight slices) with concentrations in the range of 25-50 µM (RHC80267) or 50 µM (orlistat).
The recovery during drug wash-out was accelerated by the CB1 receptor agonist HU210. A wash-in of HU210 (10 µM) for 2 min led to an almost complete recovery of the responses (Figure 17 A, B, green traces). The described effects are very similar to those induced by blockage of CB1 receptors as published previously (Czesnik et al., 2007, Figure 17 C).

3.2.3 Differential expression of the DAGL isoforms within the OE

While the above data demonstrate that the suppression of odorant responses was brought about by the endocannabinoid 2-AG, produced by a DAGL, the production site of 2-AG, i.e. the site of DAGL activity remained unclear so far. Therefore the expression of the DAGL in the OE, specifically the expression of the α and β isoforms was localized. Olfactory receptor neurons and sustentacular cells, which could easily be distinguished on the basis of their characteristic morphology, were first patch-

Figure 16: Endocannabinoid system in the OE of tadpoles. PCR products for mRNAs of the CB1 receptor (lane 1), the two DAGL isoforms α and β (lane 2 and 3), MAGL (lane 4), NAPE-PLD (lane 5), and FAAH (lane 6) were electrophoretically separated in an ethidium bromide containing agarose gel and visualized with UV-light.
clamped and physiologically identified. Then the cytoplasm of the patch-clamped cell was harvested into the patch pipette for further PCR analysis. The mRNA of olfactory marker protein 1 (OMP1) and of cytokeratin type II (CYTII) were used as markers to confirm the identity of ORNs and sustentacular cells, respectively (Rössler et al., 1998; Hassenklöver et al., 2008). Five out of ten ORNs (OMP1-positive) expressed DAGLβ, and none of them expressed DAGLα. On the other hand, five out of eight sustentacular cells (CYTII-positive) expressed DAGLα, and none of them DAGLβ. In summary, 2-AG is synthesized in both ORNs and sustentacular cells, though by two different isoforms of the DAGL (see examples in Figure 18). The β-isoform is active in ORNs and the α-isoform in sustentacular cells.
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3.2.4 DAGLα expression is enhanced after food-deprivation

In order to find a functional link between the nutritional or hunger state of an animal on the one hand and 2-AG synthesis on the other, it was investigated whether hunger affected the expression of DAGL. To this end mRNA of both DAGL isoforms were obtained and analyzed from five groups of animals using real-time PCR. The first and second group of animals were food-deprived either for 6 h (group A_{6h}) or for 12 h (group A_{12h}) before analyzing their mRNA levels. The third and forth group were fed to satiety for 2 h after having been food-deprived for 6 h (group B_{6h}) or 12 h (group B_{12h}). A control group of animals (group C) was fed to satiety for 2 h before measurements. The mRNA expression levels for the A- and B-groups were normalized to those of the control group (Figure 19, grey line).

Comparing the expression levels of DAGLα (blue; sustentacular cells) and DAGLβ (red; ORNs), hunger clearly had no effect on 2-AG production in ORNs (Figure 19, red points), since the normalized changes of the DAGLβ (ORNs) by hunger (groups A_{6h} and A_{12h}) or refeeding after hunger (groups B_{6h} and B_{12h}) were

Figure 18: Differential DAGLβ and α expression in ORNs and sustentacular cells. Single-cell RT-PCR revealed mRNA of DAGLβ solely in ORNs (OMP1-positive cells) and mRNA of DAGLα in sustentacular cells (CYTII-positive cells).
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Figure 19: DAGL mRNA expression is regulated upon food-deprivation. Relative expression levels (i.e. normalized to the control group) of DAGLα in sustentacular cells (blue points) and β in ORNs (red points) in the OE exposed to four nutritious states. DAGLα expression levels were affected by the various nutritious conditions (group A₆h, food-deprived for 6 h, n = 7, 1.45 fold; group B₆h, food-deprived for 6 h and refed for 2 h, n = 7, 1.32 fold; group A₁₂h, food-deprived for 12 h, n = 7, 1.50 fold; group B₁₂h, food-deprived for 12 h and refed for 2 h, n = 7, 1.52 fold). DAGLβ expression levels were not affected by the various nutritious conditions (0.99 fold, 1.07 fold, 0.97 fold, 1.05 fold (n = 7), respectively).

0.99, 1.07, 0.97, and 1.05, respectively (n = 7). In contrast, in sustentacular cells, DAGLα expression was significantly enhanced after food deprivation for both 6 h (group A₆h) and 12 h (group A₁₂h). On average the mRNA expression levels were 1.45 times (n = 7; A₆h) or 1.50 times (n = 7; group A₁₂h) higher than in the control group. Refeeding for 2 h after 6 h food deprivation diminished the increase slightly (mean: 1.32 fold; n = 7; group B₆h), while the enhanced expression levels after 12 h food deprivation showed no recovery (mean: 1.52 fold; n = 7, group B₁₂h).

3.2.5 The endocannabinoid level tunes odor thresholds of individual ORNs

The above data suggested that 2-AG modulates the sensitivity of ORNs. As to possible sensitivity measures, the obvious candidates were the concentration at
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The dose-response curve is half-maximum, $K_{1/2}$, or the threshold concentration below which an ORN shows no response to the stimulus, $c_{th}$. Dose-response curves of ORNs for a number of stimuli (arginine, methionine and lysine) were measured (done by B. Gutermann) and fitted to a Boltzmann function. Figure 20 A, B shows

**Figure 20: Dose-response relationships of $[\text{Ca}^{2+}]_i$ transients induced by single amino acids.** (A) $[\text{Ca}^{2+}]_i$, transients of a single ORNs elicited by increasing concentrations (0, 0.2, 1, 2, 10, 20, 50, 100, 200, 500, 1000 and 2000, in µM) of lysine (lys) are shown. The detection threshold concentration (*) in this example is 1 µM. (B) the amplitudes obtained by the demonstrated dose-response-measurements were fitted by a Boltzmann equation. (C) Histograms of all investigated ORNs were classified for the three amino acids lysine (lys), arginine (arg) and methionine (met) according to their individual detection thresholds. The black lines indicate the application of the odorants.
ORN responses to lysine together with the corresponding dose-response curve as an example. The midpoint slopes of the 65 dose-response curves measured varied considerably (by a factor > 10) so that curves having the same $K_{1/2}$ had quite different $c_{th}$ values (not shown). Therefore the threshold concentration, $c_{th}$, was preferred as a convenient measure of sensitivity, whereby $c_{th}$ is defined as the concentration below which, under control conditions, no responses could be measured. Specifically, the first data point of the monotonic increase of the dose-response-curve was taken as the detection threshold $c_{th}$. Note that this definition refers to control conditions (i.e., no food shortage and no drugs applied).

The detection thresholds varied from ORN to ORN over a wide range. Figure 20 C gives the threshold distributions for the three odorants used.

To investigate the effect of 2-AG on the odorant detection threshold of a specific ORN, a control experiment as shown in Figure 21 A was carried out first. The orange trace gives an arginine-induced $[Ca^{2+}]_i$ transient at the detection threshold $c_{th}$ (in this case, 20 µM). Expectedly, a higher odorant concentration induced a larger response amplitude and a shorter response latency (black trace, 50 µM), while concentrations below $c_{th}$ failed to elicit a response in this ORN (blue trace, 10 µM). Importantly, this response behavior was well reproducible (Figure 21 B, orange and brown traces, blue and light blue traces). Now the slice was superfused with the DAGL blocker RHC80267, which consistently led to response failures at $c_{th}$ (Figure 21 C, red trace, RHC80267, 50 µM), meaning that the threshold $c_{th}$, under the experimental condition of less 2-AG being produced was shifted to a higher value, i.e., $c_{th} > c_{th}$. Mimicking the presence of 2-AG by wash-in of the CB1 receptor agonist HU210 (10 µM) was able to rescue the odorant responses at $c_{th}$ (Figure 21 C, green trace). Moreover, HU210 was not only able to rescue the response; frequently it also lowered the threshold so that responses could be recorded at subthreshold odorant concentrations ($c < c_{th}$). This is shown in Figure 21 D, where an odor response failure at 10 µM ($c < c_{th}$, blue trace) is transformed into a clear response at the same concentration after HU210 was added to the bath (Figure 21 D, green trace). There is thus no doubt that the sensitivities of ORNs are modulated by endocannabinoids.

While Figure 21 C gave a typical example, Figure 22 A summarizes the data for all ORNs recorded under this condition. The cells are grouped with respect to their threshold concentration $c_{th}$ (abscissa). The left (orange) column of each column
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triplet of the histogram gives the number of ORNs found to have the threshold concentration indicated on the abscissa. The middle column (red) gives the number of ORNs that show a response suppression (threshold increase) upon DAGL blockage, and the right column of each column triplet (green) shows how many ORNs regained an odor response after adding HU210 to the bath. An increase of threshold concentration upon application of RHC80267 or orlistat was observed in 54 out of 54 ORNs (52 slices; 18 cells for arginine, 21 cells for lysine and 15 cells for orlistat).

Figure 21: ORNs have individual and tunable odorant detection thresholds according to the 2-AG level in the OE. (A) ORN $[\text{Ca}^{2+}]_i$ responses to various concentrations of arg (10, 20, 50 µM). (B) The responses to the detection threshold $c_{\text{th}}$ (20 µM, orange and brown traces) and concentrations below $c_{\text{th}}$ (10 µM, blue and light blue traces) were highly reproducible. (C) After addition of RHC80267 (50 µM) to the bath solution the $[\text{Ca}^{2+}]_i$ transients induced at $c_{\text{th}}$ were abolished (red trace). Wash-out of RHC80267 with HU210 in the bath solution (2 min, 10 µM, green trace) accelerated recovery. (D) Lacking odorant response under control conditions (arg, blue trace) and reappearing of odorant response after addition of HU210 (2 min, green trace) to the bath. The black lines indicate the application of the odorants.
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Figure 22: Summary of detection threshold modulation. (A) Histogram of a group of ORNs responsive to arginine, lysine or methionine classified according to their individual detection thresholds under control conditions (orange bars). In all of these ORNs RHC80267 or orlistat led to a reduction of the response (red bars) whereby a recovery (drug wash-in of HU210 10 µM, green bars) could be observed in 42 out of 53 ORNs. (B) Histogram of groups of arginine-, lysine- or methionine-responsive ORNs (blue bars) plotted over the highest concentration where no response could be recorded. In 19 out of 38 ORNs HU210 (10 µM) permitted responses at the respective concentrations (green bars).

methionine; red bars), and the agonist HU210 led to a recovery in 42 out of these 54 ORNs (green bars).
Figure 23: PMCA but not NCX mediates CB1 receptor-induced effects. (A) Aamix-evoked [Ca^{2+}]_{i} transients of individual ORNs (black traces) were reduced and delayed after wash-in of the PMCA inhibitor CE (5 µM; red trace). After drug wash-out with bath solution, the [Ca^{2+}]_{i} transients recovered (blue traces). (B) Aamix-evoked [Ca^{2+}]_{i} transients of individual ORNs (black traces) were only reduced after substitution of Na^{+} with NMDG (NMDG25: dark grey; NMDG10: middle grey; NMDG0: light grey). After drug wash-out with bath solution, the [Ca^{2+}]_{i} transients recovered (blue traces). (C) CE-modulated [Ca^{2+}]_{i}
Figure 22 B summarizes the experiments where stimulation was at subthreshold concentrations, i.e., at $c < c_{th}$, where no responses could be elicited, and repeated the stimulation with the CB1 agonist HU210 added to the bath. With HU210 in the bath, responses were observed in 19 (green) out of 38 (blue) cells (38 slices). There was no correlation between the occurrence of this effect and the corresponding threshold concentration.

3.2.6 Preliminary: PMCA is effector molecule of the CB1 receptor in ORNs

Although the cellular effect of CB1 receptor blockage and activation was demonstrated, the underlying signaling systems are unknown so far. CB1 receptor signaling mostly modulates intracellular Ca$^{2+}$ levels. Ca$^{2+}$ is extruded from ORNs by the plasma membrane Ca$^{2+}$ ATPase (PMCA; Lischka and Schild, 1993; Castillo et al., 2007) and the Na$^+$-Ca$^{2+}$-exchanger (NCX; Reisert and Matthews, 1998; Lucero et al., 2000).
First, a possible involvement of PMCA in CB1 signaling was investigated. The PMCA blocker carboxyeosin (CE) was used during odorant stimulation of ORNs with amino acids. The black trace in Figure 23 A shows a \([\text{Ca}^{2+}]\) transient upon stimulation with amino acids under control conditions. Wash-in of CE (5 µM; 10 min; Figure 23 A, red trace) delayed the response and reduced its amplitude. This effect is similar to that observed for blockage of 2-AG synthesis (Figure 17). A subsequent wash-out of the drug recovered the response (Figure 23 A, blue trace; observed in 28 out of 28 cells in three slices).

In the next step, the involvement of NCX in CB1 signaling was examined. To achieve a stop or even a reverse mode of the NCX and thus, retain \(\text{Ca}^{2+}\) ions in the cell, extracellular \(\text{Na}^+\) was substituted for NMDG (NMDG25: 25 mM NaCl, NMDG10: 10 mM NaCl, NMDG0: 0 mM NaCl; Schild et al., 1994). These conditions reduced the amplitude of \([\text{Ca}^{2+}]\) responses (Figure 23 B, grey traces) compared to the control (Figure 23 B, black trace). Restorage of extracellular \(\text{Na}^+\) to 98 mM recovered the response (Figure 23 B, blue trace). A delay of the response could not be observed by NCX interference (observed in 28 out of 28 cells in four slices).

In the following experiments it was investigated whether these described effects occur downstream of the CB1 receptor or whether they are mediated CB1 receptor-independent. Thus, the experiment was slightly modified and the altered \([\text{Ca}^{2+}]\) responses upon PMCA or NCX blockage were tried to be rescued by wash-in of the CB1 receptor agonist HU210. The black traces in Figure 23 C and D are \([\text{Ca}^{2+}]\) responses to amino acids under control conditions. Wash-in of CE (5 µM; Figure 23 C, red trace) reduced and delayed the response and substitution of extracellular \(\text{Na}^+\) by NMGD (in this case NMDG 10; Figure 23 D, grey trace) only reduced the response. Subsequent wash-in of HU210 (20 µM) rescued the response upon CE (Figure 23 C, green trace; observed in 21 out of 26 cells in four slices), but had no effect on the responses altered by NMDG10 (HU210, 10 µM; Figure 23 D, green trace; observed in 52 out of 52 cells in four slices). This suggests that PMCA is located downstream in the CB1 receptor signaling cascade in ORNs, but not NCX.

To further verify this finding, it was tested, whether CE still triggers the described effect when the NCX is out of action. The control response to amino acid application in Figure 23 E (Figure 23 E, black trace) was reduced in amplitude upon NMDG10 (Figure 23 E, grey trace). Wash-in of CE (5 µM; Figure 23 E, green trace)
3.2 Endocannabinoid modulation in the olfactory epithelium

further reduced and additionally delayed the $[\text{Ca}^{2+}]$ response. Wash-out with bath solution recovered the transient (Figure 23 E, blue trace; observed in 45 of 45 cells in three slices).

To localize the CB1/PMCA-mediated $[\text{Ca}^{2+}]$ change in ORNs, single dendrites were imaged during CE incubation of tissue slices. Exemplarily, a fluo-4-loaded cell is shown in Figure 24 (left side) as a z-projection. Regions of interest were arranged as indicated by the numbers. The $[\text{Ca}^{2+}]$ levels in these regions upon CE in the bath solution was normalized to those obtained during control conditions (five images accessed prior to the experiment, grey line). $[\text{Ca}^{2+}]$ increased in all regions 2.5 min after CE wash-in, the increase was highest at the knob region (red trace) and smallest close to the soma (black region). Similar results were obtained for three cells. First experiments with the CB1 antagonist AM281 revealed similar results (data

![Figure 24: $[\text{Ca}^{2+}]$ increases in the distal dendrite upon CE. Left side: fluo-4-loaded ORN with indicated regions of interest. Right: The $[\text{Ca}^{2+}]$ level upon CE was normalized to these of control conditions (grey line). $[\text{Ca}^{2+}]$ increased in all regions 2.5 min after CE wash-in. The increase was highest at the knob region (red trace) and smallest close to the soma (black region). Scale bars: 1 min and $\Delta F/F = 250\%$.](image-url)
Xenopus laevis
Rattus norvegicus
Homo sapiens

CB1

DAGLα

DAGLβ

NAPE-PLD

FAAH

MAGL

CB2

GPR55

Xenopus laevis
Rattus norvegicus
Homo sapiens

Dopamine receptor 2

Dopamine receptor 2B

Adiponectin receptor 1

Adiponectin receptor 2

Orexin receptor 1

Orexin receptor 2

Leptin receptor

GPR55
not shown; n = 3). However, more experiments have to be carried out to draw a final conclusion.

3.2.7 Endocannabinoid system and receptors for other modulators in the OE of various species

*Xenopus laevis* tadpoles served as an animal model in all experiments so far. Of course, it is interesting to find out, if the endocannabinoid system is located also in mammals, especially in humans. Therefore, the mRNA in OEs of rats and a human was analyzed in addition to mRNA of tadpoles.

First, it was examined, whether mRNA for the synthesis and degradation enzymes for 2-AG (DAGLα, DAGLβ, FAAH) and anandamide (NAPE-PLD, MAGL) are present in the OE (Table 6). In fact, PCR products for the corresponding mRNAs were detected in all samples investigated. CB1 receptor mRNA was detected in the OE of tadpoles and rats, but not in the human sample. In rats, CB2 receptor and GPR55 mRNA was additionally found. In humans, only GPR55 mRNA was detected as a receptor for endocannabinoids.

Since a link of the nutritive status to the endocannabinoid system in the OE was shown in this thesis (Figure 19), the presence of several receptors, among them receptors for orexigenic substances was investigated on the transcriptional level in OEs (Table 6). In tadpoles, the mRNA for dopamine, adiponectin, and leptin receptors were localized to the OE. Rats exhibited mRNA for dopamine, adiponectin, orexin, and leptin receptors in the OE and the human sample showed adiponectin and leptin receptor mRNA, but not dopamine and orexin receptor mRNA.

Table 6: mRNA of the components of the endocannabinoid system and receptors for other modulatory substances. In this table all investigated mRNAs of OEs of tadpoles (n = 3), rats (n = 3), and a human (n = 1) are shown. Components of the endocannabinoid system (CB1, DAGLα, DAGLβ, NAPE-PLD, FAAH, MAGL, CB2, GPR55) and receptors for orexigenic substances (dopamine receptors A and B, adiponectin receptors 1 and 2, orexin receptors 1 and 2, leptin receptor) are listed. Empty table elements mirror mRNA, for which the gene sequence is not known or was not analyzed.
4 Discussion

4.1 The styryl dye FM1-43 permeates and blocks CNG channels in olfactory neurons of larval *Xenopus laevis*

The first question I raised in the introduction was by which mechanisms odors are encoded by the activity of ORN subsets. An essential step toward solving this issue is to establish a tool in order to differentiate ORN subsets and to specifically interfere with one ORN subset pharmacologically. Here, the action of FM1-43 in the OE was investigated and the mechanism by which it acts was characterized.

4.1.1 FM1-43 stains ORNs

FM1-43, among other styryl dyes, is mainly used in the neurosciences to monitor membrane trafficking. Since this substance reversibly stains membranes but does not pass them, and because it changes its fluorescent properties according to the hydrophobicity of its environment, FM1-43 became a powerful tool to investigate synaptic vesicle recycling and synaptic transmission (Cochilla *et al*., 1999; Kidokoro *et al*., 2004; Kay *et al*., 1999). However, there are some studies, which describe the staining of neuronal cells by styryl dyes. FM1-43 labels several sensory and neuronal cells, e.g., sensory hair cells in the lateral line organ, the cochlea of various vertebrate species (Nishikawa and Sasaki, 1996; Seiler and Nicolson, 1999; Gale *et al*., 2001; Meyers *et al*., 2003), Merkel cells, taste buds, nociceptive fibers as well as primary sensory neurons in the trigeminal (V), geniculate (VII), petrosal (IX), nodose (X), and dorsal root ganglia (Meyers *et al*., 2003; Drew and Wood, 2007; Drew *et al*., 2007).

Nishikawa and Sasaki (1996) reported that FM1-43 labeled epidermal cells at nasal pits. The labeled cells were not further identified. Three years later, FM1-43 was shown to label dissociated ORNs (Rankin *et al*., 1999). In these experiments FM1-43 was internalized and appeared in the cell body, dendrite, and knob after
stimulation with L-glutamate, but the dye could not be located to the cytosol or the plasma membrane due to technical reasons. Rankin et al. (1999) postulated a novel endocytosis-like mechanism for the dye uptake. In the present work, I found no evidence for this hypothesis. FM1-43 was shown herein to stain ORNs in the OE of Xenopus laevis tadpoles when living animals were incubated in a solution of distilled water and the dye (Figure 9). Fluorescence was observed in the cytosol and was absent only in the nucleus. The same cytosolic staining pattern as in this thesis was also observed for hair cells by Gale et al. (2001) and Meyers et al. (2003).

### 4.1.2 FM1-43 uptake in ORNs through CNG channels

Uptake of dyes through plasma membrane channels seems to be a more general process than previously assumed. For example, YO-PRO permeates purinergic receptors (Khakh et al., 1999) and TAE permeates the mechanoelectric transduction channel. Besides hair cell mechanotransducer channels, other sensory channels like the vanilloid receptor TRPV1, the purinergic receptor P2X\(_2\) and mechanoelectric transduction channel of dorsal root ganglion cells (Meyers et al., 2003; Drew and Wood, 2007) were shown to be permeable for FM1-43.

Meyers and coworkers (2003) compared FM1-43 staining of hair cells with that of FM3-25, a structurally related styryl dye with similar properties. FM3-25 only labeled the plasma membrane of hair cells whereas FM1-43 caused a cytosolic staining. This comparison shows that FM1-43 stains certain cells by a different mechanism than membrane insertion, namely channel permeation.

For FM1-43 staining of ORNs, living tadpoles were incubated in a solution of the dye and distilled water. The cells at the apical side of the OE are connected by tight junctions that prevent the diffusion or the transport of molecules from the principal cavity into the tissue (Miragall et al., 1994; Steinke et al., 2008). Therefore, dye uptake certainly occurred through the plasma membrane exposed to the principal cavity: cilia, microvilli, and/or dendritic knobs come into consideration. Indeed, FM1-43 uptake in hair cells also occurred at the stereocilia (Gale et al., 2001; Meyers et al., 2003), where removal of the cilia prevented dye uptake.
Since FM1-43 fluorescence was cytosolic, since it built up rapidly, and since CNG channels are located on cilia, it was checked whether CNG channels are permeable for FM1-43. The well-known permeability of divalent ions through CNG channels as well as the effect of two non-specific blockers of CNG channels were used here. FM1-43-staining of ORNs was blocked when divalent ions were present during dye internalization (Figure 11). Similarly, FM1-43 competes with other cations for uptake through the mechanoelectric transducer channel in hair cells (Nishikawa and Sasaki, 1996; Seiler and Nicolson, 1999; Gale et al., 2001).

Furthermore, the unspecific CNG channel blockers LY-83,583 and amiloride prevented dye uptake in ORNs as presented in this thesis (Figure 12). Endocytosis-independent FM1-43 uptake was also investigated by Meyers et al. (2003) in bullfrog and mice hair cells. In that study cells rapidly filled by diffusion of FM1-43 from the apical to the basal pole. Mechanical closure of the mechanotransducer channel was sufficient to block dye uptake. Gale and coworkers obtained similar results in 2001. Additionally, they demonstrated that hair cells of Myo 7a mutants cannot be labeled with FM1-43. In these mutants, the mechanoelectric transduction channels, and thus the gates for FM1-43 internalization, are closed at rest. In addition, treatment of hair cells with the Ca²⁺ chelator EGTA, a condition which breaks tip links and thus closes the mechanoelectric transduction channel, abolished subsequent dye loading.

In this study, FM1-43 entered ORNs in the absence of an externally applied stimulus. Generally, the exclusion of any kind of stimulation can hardly ever be met in olfactometry. Apart from this caveat, CNG channels in ORNs are reported to gate spontaneously and ligand-independent, thereby producing a detectable macroscopic conductance (Kaupp and Seifert, 2002). Kleene (2000) estimated the open probability of CNG channels due to spontaneous gating in dissociated grass frog ORNs to be approximately 0.03. Tibbs and coworkers (1997) calculated an open probability of 0.002. Their model was an exogenous expression system with the α subunit of the catfish olfactory CNG channel. Combined with the incubation time of several minutes, this would allow spontaneous dye uptake in ORNs. This concept also holds true for hair cells: these can also be loaded with FM1-43 in the absence of a stimulus. The open probability of the mechanoelectric transduction channel at rest is 0.1 to 0.2. A stimulus essentially increases the channel open probability and the current across the membrane (Grant and Fuchs, 2007).
4.1.3 Only ORNs endowed with the cAMP transduction machinery internalize FM1-43

FM1-43 always stained a subset of ORNs in the OE. The identity of this subset was characterized herein by three experiments: First, FM1-43 staining of the ORN subset was blocked when divalent ions were present during dye internalization (Figure 11). Second, unspecific CNG channel blockers inhibited dye internalization (Figure 12). Third, FM1-43-stained ORNs exhibited the cAMP-dependent transduction cascade because they could be stimulated with forskolin and uncaging of cAMP (Figure 13).

These findings indicate that FM1-43 permeates CNG channels and stains ORNs with the cAMP-dependent transduction cascade. In a number of publications it has been reported that only a fraction of *Xenopus laevis* ORNs possess the canonical, cAMP-dependent olfactory transduction cascade (Manzini *et al.*, 2002; Manzini *et al.*, 2003b; Czesnik *et al.*, 2006). Other ORNs in the OE detect odorants, e.g. amino acids, via a cAMP-independent transduction mechanism. FM1-43 must thus be supposed to stain cAMP-dependent ORNs when permeating CNG channels. Olfactory receptor neurons that could not be stained with FM1-43 are therefore believed to express a different kind of generator channel. If FM1-43 permeated those channels too, the vast majority of ORNs in the OE would be stained. As this was not the case, it can be concluded that the ORN generator channels involved in the detection of amino acids are not permeable for FM1-43.

In studies, in which hair cells were labeled with FM1-43, the staining was observed in all cells exposed to the dye (e.g. Seiler and Nicolson, 1999; Meyers *et al.*, 2003). This may be due to the fact that the transduction mechanism in hair cells seems to be identical throughout the hair cell population in the cochlea. The identity of the mechanoelectric transduction channel is still unknown. However, it is a nonselective cation pore, permeable for K⁺ and Ca²⁺ (Grant and Fuchs, 2007; Phillips *et al.*, 2008). FM1-43 is thus used as a marker for mature hair cells (Doyle *et al.*, 2007).
4.1.4 Extracellular FM1-43 inhibits cation currents through CNG channels

To date, little systematic effort was made to develop potent and specific pharmacological agents that inhibit CNG channels. CNG channel blockers known so far are unspecific and have to be used in high concentrations causing many side effects. Others block channels only at positive membrane potentials. FM1-43 inhibits CNG channels at the resting membrane potential at low concentration (Figure 14). In the following, I will provide a short overview of CNG channel blockers with their properties concerning potency and specificity. Then, I will discuss FM1-43 as a CNG channel blocker:

L-cis-diltiazem is probably the CNG channel blocker, which has been studied most extensively (Koch and Kaupp, 1885; Haynes, 1992; Brown et al., 2006). It inhibits the rod photoreceptor CNG channels noncompetitively and in a voltage-dependent manner. The $K_{1/2}$ of L-cis-diltiazem is in the low micromolar range at +30 mV and increases with negative voltages. This blocker affects CNG channels from the cytoplasmatic side, and it is non-specific. Amiloride is often used as an epithelial $Na^+$ channel blocker at low micromolar concentrations, and it also blocks rod and olfactory CNG channels (used in Figure 12; Frings et al., 1992; Brown et al., 2006). The inhibition is strongly voltage-dependent and least effective at the resting membrane potential. D-600 and verapamil, two amiloride derivates, have similar properties (Frings et al., 1992). Dichlorobenzamil, another derivate of amiloride has more promising characteristics (Nicol et al., 1987). It blocks CNG channels at low micromolar concentrations, and it is relatively voltage-independent. Nevertheless, amiloride derivates cannot be considered as selective antagonists since they inhibit also the $Na^+$-$Ca^{2+}$ exchanger and voltage-gated $Na^+$, $Ca^{2+}$ and $K^+$ channels in the similar concentration range. LY-83,583 is another unspecific CNG channel blocker. It was used in this thesis at a high concentration (Figure 12). Besides this function, it blocks cGMP production, inhibits intracellular $Ca^{2+}$ release, and blocks the effects of nitric oxide (Leinders-Zufall and Zufall, 1995). Tetracaine, a local anesthetic, blocks $Na^+$, $Ca^{2+}$ and CNG channels at micromolar concentrations. The block of CNG channels by tetracaine is voltage- and state-dependent. Two studies of Karpen’s lab (Ghatpande et al., 2003; Strassmaier et al., 2005) reported on tetracaine analogues with less side effects. The most potent inhibitor for monomeric CNG channels is
4.1 FM1-43 permeates and blocks CNG channels

Pseudochitoxin (Brown et al., 1999; Kaupp and Seifert, 2002). The $K_{1/2}$ is 5 to 100 nM. Concerning heteromeric channels, the pseudochitoxin blocks CNG channels several odors of magnitudes less effectively.

Because of these unfavorable properties of the blockers, the concentrations of LY-83,583 and amiloride were very high (200 µM and 1 mM, respectively) in the presented experiment in Figure 12. Nevertheless, staining of ORNs with FM1-43 was abolished when these blockers were present.

In contrast to many CNG channel blockers mentioned above, FM1-43 blocks at resting membrane potentials. Cells can be stained in vivo without stimulation. Furthermore, 10 µM FM1-43 reduced the CNG current to ~25 %. This was measured in the absence of $Ca^{2+}$ and $Mg^{2+}$ at the resting membrane potential. The blockage occurred from the extracellular site as proven in Figure 15.

FM1-43 was also described as a blocker of cation currents in two other studies: Gale et al. (2001) observed that extracellular FM1-43 reversibly blocks mechanotransduction in cochlear hair cells in culture. FM1-43 reduces the currents in a voltage-dependent way. The block is most effective at -4 mV ($K_d = 1.2 \mu M$) and less effective at large positive and negative potentials. Furthermore, the block is strongly dependent on extracellular $Ca^{2+}$ and most effective at low $Ca^{2+}$ concentrations. In a study by Drew and Wood (2007) extracellular FM1-43 blocked rapidly- and slowly-adapting mechanically activated cation currents in cultured dorsal root ganglion neurons. The $K_d$ is 5 µM and 3 µM, respectively. The block was equally efficient at voltages of -70 and -35 mV, but it was significantly reduced at positive holding potentials. At low extracellular $Ca^{2+}$ concentrations the FM1-43 block of the currents was more effective.

With this knowledge, one of the first experiments I presented, i.e. the determination of odor sensitivity of FM1-43-loaded ORNs (Figure 10), becomes more comprehensible. Only few FM1-43-stained ORNs could be stimulated with an odor-mixture. The stimuli for these cells included amines and alcohols and elicited only very small $[Ca^{2+}]_i$ transients. A second stimulation with the odors led to a reduced response. Considering these aspects, one can conclude that while permeating CNG channels FM1-43 blocks the ionic current through these channels. In this way odorant responses are prohibited. This is very useful as there are virtually no other potent and specific blockers for CNG channels known.
Figure 25: Action of FM1-43 in the OE. FM1-43 labels the ORN subset endowed with the cAMP-dependent transduction cascade (right side, ORN highlighted in red). Extracellular FM1-43 inhibits CNG channels. ORNs that use a different transduction mechanism and have a different generator channel are not stained with FM1-43 (left side, ORN in grey). PC, principal cavity; ON, olfactory nerve; od, odorant, SC, sustentacular cell.
4.1.5 Conclusion

Taken together, FM1-43 appears to exert a permeation block of CNG channels (Figure 25). It is a novel mechanism to label a distinct subset of ORNs, and conversely, to identify non-labeled cells such as sustentacular cells or ORNs that don't use cAMP in their transduction cascade. Further it allows staining and blocking \textit{in vivo} and under physiological conditions. It seems therefore particularly useful for studies of olfactory transduction cascades. Finally the fluorescence of FM1-43 may turn out to be well-suited for studying ciliary processes and channel densities.
4.2 Modulation of processing in olfactory receptor neurons by the endogenous cannabinoid system

In the introduction, I pointed out that ORNs cannot be considered as static units. The physiological state of the organism may influence the fine-tuning of odor detection. Since many modulatory systems in the OE seem to work in parallel, one has to analyze all of these systems separately, and then investigate their interplay. Here I examined the effect of the endocannabinoid 2-AG on olfactory processing. In the following sections, I present the functional meaning of endocannabinoid modulation in the OE, its action, and its mechanism of action.

4.2.1 The endocannabinoid 2-AG acts in the OE

Several studies showed that CB1 receptors or the related mRNA can be found at different stages of the central olfactory system (Cesa et al., 2001; Egertová and Elphick, 2000). A study of Migliarini et al. (2006) demonstrated the presence of CB1 receptor mRNA in the OE of *Xenopus laevis* tadpoles at stage 46. Recently, Czesnik and coworkers (2007) revealed that ORNs are modulated by cannabinoids. CB1 receptor antagonists reduced and delayed odor-induced responses of ORNs, and the CB1 receptor agonist HU210 accelerated the recovery of these responses. Additionally, they localized CB1 receptors on ORN dendrites in the OE. These results indicate the presence of the endogenous cannabinoid system in the OE. However, the type of endocannabinoid acting in the OE was not described. Therefore, an mRNA analysis of the components of the endocannabinoid system in the OE of tadpoles was performed. Indeed, mRNA of the CB1 receptor as well as the synthesis and degradation enzymes for 2-AG (DAGLα, DAGLβ, MAGL) and anandamide (NAPE-PLD, FAAH) were detected (Figure 16). This implies that both 2-AG and anandamide are synthesized in the OE.

2-AG is more abundant in the brain than anandamide (Sugiura et al., 2006). For this reason, its action was investigated by blockage of 2-AG synthesis with the DAGL inhibitors RHC80267 or orlistat (Hashimotodani et al., 2008) in the present
work. This decreased and delayed the odor-induced $[\text{Ca}^{2+}]_i$ transients. Wash-in of the CB1 receptor agonist HU210 rescued the responses (Figure 17). The same effects were obtained with the CB1 receptor antagonists AM281, AM251, and LY320135 by Czesnik and coworkers (2007).

Besides the olfactory system, other sensory systems are also known to be modulated by the endocannabinoid system. For example, CB1 receptors are located on photoreceptors and bipolar cells in the visual system. There, cannabinoids speed up the dynamics of the phototransduction deactivation cascade in cones (Straiker et al., 1999; Struik et al., 2006). In addition, CB1 receptors are expressed on dorsal root ganglion cells, and may play a role in the spinal nociceptive system (Morisset et al., 2001). The endocannabinoid modulation of sensory output at the most peripheral stage may thus be a common feature of these sensory systems.

### 4.2.2 Cellular localization of 2-AG synthesis and its functional meaning

2-AG acts in the OE, and it is synthesized by DAGL$\alpha$ and $\beta$. However, the cellular localization of 2-AG synthesis was not known so far. In this thesis, mRNA of both the DAGL$\alpha$ and $\beta$ isoforms were detected by single-cell PCR in sustentacular cells and ORNs, respectively (Figure 18). In ORNs appears to exist an autocrine pathway since 2-AG is produced by DAGL$\beta$ in ORNs, and since it acts on CB1 receptors on ORNs (Czesnik et al., 2007). In contrast, DAGL$\alpha$ mRNA is solely expressed in sustentacular cells, indicating an additional paracrine route of 2-AG action in the OE.

2-AG production by DAGL$\alpha$ in sustentacular cells is enhanced upon food-deprivation

2-AG is produced by DAGL$\alpha$ in sustentacular cells and acts paracrine on ORN dendrites. Sustentacular cells insulate ORNs (Breipohl et al., 1974; Getchell and Getchell, 1992; Farbman, 1992; Morrison and Moran, 1995) and regulate mucus secretion and ion homeostasis of the extracellular compartment (Getchell and Getchell, 1992; Hansen et al., 1998). In this thesis, a novel role of sustentacular cells is indicated. 2-AG is secreted by sustentacular cells and modulates the activity of ORNs. Additionally, DAGL$\alpha$ mRNA expression in the OE was found to be enhanced after food deprivation, whereas DAGL$\beta$ expression was not affected by the various
nutritious conditions (Figure 19). This allows to conclude that 2-AG production in sustentacular cells by DAGLα is enhanced by hunger and acts via a paracrine route upon CB1 receptors on ORN dendrites.

The endocannabinoid system is known to play a crucial role in food intake and energy homeostasis (Aimé et al., 2007). For instance, in the teleost fish Carassius auratus (Soderstrom et al., 2004), in the zebra finch (Kirkham et al., 2002), and in rodents (Di Marzo et al., 2001; McLaughlin et al., 2003), brain endocannabinoids seem to act as orexigenic mediators. In addition, AM251 induces suppression of rat food intake and food-reinforced behavior in rats (Mousley et al., 2006). The link between exocannabinoids and increased food intake is well-known (Hart et al., 2002; Verty et al., 2005). A previous study has shown that CB1 receptor antagonists diminish and delay odor responses (Czesnik et al., 2007). In this thesis, a functional link between 2-AG as a modulator in the OE and the nutritious state of an animal is shown. It was demonstrated that there is an endocannabinoid-system-mediated crosstalk between the neuronal control of feeding, e.g. olfaction, and the nutritional state.

A similar concept was suggested for orexin and leptin, which are hormones involved in energy metabolism. Orexins are synthesized within the OE. Orexin receptors were localized among others on the apical part and microvilli of sustentacular cells and knobs and cilia of ORNs in the OE. Thus, a possible modulation of olfactory perception by these neuropeptides is probable (Caillol et al., 2003). Intranasally applied orexin A restores olfactory function in narcolepsy (Baier et al., 2008), and more importantly, cerebroventricular injection of orexin results in an increased olfactory sensitivity (Julliard et al., 2007). Leptin and its receptors were also found on sustentacular cells and ORNs (Baly et al., 2007). Fasting caused a significantly enhanced transcription of both leptin and leptin receptors. Furthermore, leptin decreases odor-induced receptor potentials (Savigner et al., 2009) and sensitivity in a behavioral experiment (Julliard et al., 2007).
2-AG produced by DAGLβ in ORNs may play a role in ORN protection or differentiation

2-AG is also produced by DAGLβ in ORNs and acts autocrine on CB1 receptors on ORN dendrites. As to the autocrine pathway, no particular modulation was found. However, the following two functional meanings could be plausible:

First, odor-induced increase of $[Ca^{2+}]$i can be supposed to mediate 2-AG release (as reported in other systems, Szabo et al., 2006; Hashimotodani et al., 2007) and a subsequent increase of sensitivity and signal-to-noise ratio of responses to odors.

Second, autocrine endocannabinoid signaling appears to be a key regulatory signaling network for the wiring of the brain during development (Harkany et al., 2008), and may play a role in the constantly regenerating ORNs (Schwob, 2002) in the OE. Interference with the endocannabinoid system using pharmacological inhibitors disturbs axon pathfinding and fasciculation (Watson et al., 2008). Williams and coworkers showed that DAGL-dependent activation of neuronal CB1 receptors and CB1 agonists stimulate the growth of axons while CB1 antagonists inhibit this process (Williams et al., 2003). For instance, developing pyramidal cells rely on endocannabinoid signaling to initiate the elongation and fasciculation of their long-distance axons (Mulder et al., 2008).

In the future, selective blockers will allow to study the effect of autocrine acting 2-AG, which is produced by DAGLβ.

4.2.3 2-AG level modulates odor detection thresholds

Recently, several studies dealing with the influence of the nutritious state on the neurophysiology of olfactory information processing have been published. In these studies an altered sensitivity of ORNs could indirectly be attributed to the effects of modulators like neuropeptide Y, leptin or orexin (Mousley et al., 2006; Caillol et al. 2003; Getchell et al., 2006). Here, detection thresholds of arginine, lysine, and methionine were investigated. Their dose-response relationships and detection thresholds at cellular resolution were described using confocal fluo-4 Ca$^{2+}$-imaging (Figure 20). The findings show that response thresholds are distributed over a
distinct concentration range between 0.2 and 200 µM, which has also been described by Duchamp-Viret and coworkers for ORNs in rat and adult frog (Duchamp-Viret et al., 2000). The classical view is that odorant detection thresholds are determined by the affinity and expression level of olfactory receptors (Kajiya et al., 2001; Malnic et al., 1999; Saito et al., 2009), olfactory receptor dimerization (Neuhaus et al., 2004) as well as amplification and adaptation in the transduction cascade (Takeuchi and Kurahashi, 2008). Herein, the significant contribution of endocannabinoids to the control of odorant thresholds is shown (Figure 21, Figure 22). These findings support the view that 2-AG acts as an orexigenic modulator in the OE by increasing and decreasing the sensitivity of ORNs to odorants during phases of hunger or satiety. As a consequence, the concept of a “threshold as a well defined and constant concentration below which an ORN does not respond to a specific odorant” is no longer applicable. The threshold under control conditions, $c_{th}$, as it has been used herein, may serve as a simple and convenient definition, but it should only be used if the conditions are sufficiently well-defined.

4.2.4 Novel CB1 transduction cascade in ORNs: PMCA as effector molecule

The preliminary results concerning the transduction mechanism of CB1 receptors suggest (Figure 23, Figure 24) that the PMCA is involved in CB1 signaling. CB1 receptor activation would enhance PMCA activity and thus decrease $[\text{Ca}^{2+}]_i$, whereas CB1 blockage would reduce PMCA activity and thus increase $[\text{Ca}^{2+}]_i$. The changes in $[\text{Ca}^{2+}]_i$ levels could be localized to the apical dendrite and knob in the experiments performed and will be verified in future experiments. It is known, that CB1 receptor activation modulates $[\text{Ca}^{2+}]_i$ (e.g. Straiker et al., 1999; Huang et al., 2001; Mackie et al., 1995). However, the PMCA as an effector molecule of the CB1 receptor in ORNs would propose a novel mechanism of CB1 receptor signaling. Certainly, the messengers in between the CB1 receptor and the PMCA still have to be identified.

Another question, which has to be solved in a future study is, how altered $[\text{Ca}^{2+}]_i$ levels in ORNs mediate the reduction and the delay of odorant-evoked $[\text{Ca}^{2+}]_i$ transients. One hypothesis is that changes in $[\text{Ca}^{2+}]_i$ induced by PMCA may affect
olfactory transduction by mimicking an adaptative state of the respective ORN to odorants and thus modulate ORN sensitivity. Three different types of adaptation for which elevated \([\text{Ca}^{2+}]_i\) is crucial, were described for ORNs. First, Ca\(^{2+}\)-calmodulin binding to the CNG channel decreases its affinity for cAMP. For this reason an equal stimulus results in smaller responses (Chen and Yau, 1994; Liu \textit{et al.}, 1994).

Second, Ca\(^{2+}\)-activated calmodulin-dependent protein kinase II inhibits the adenylate cyclase III by phosphorylation and attenuates odorant responses (Wei \textit{et al.}, 1998; Leinders-Zufall \textit{et al.}, 1999). Third, Ca\(^{2+}\)-activated calmodulin-dependent protein kinase II targets the phosphodiesterase 1C, which then probably enhances cAMP destruction (Yan \textit{et al.}, 1995; Borisy \textit{et al.}, 1992).

Further experiments will elucidate the CB1 transduction cascade in ORNs and the effect of CB1-receptor induced \([\text{Ca}^{2+}]_i\) changes by PMCA in ORNs.

4.2.5 Endocannabinoid and other modulatory systems exist in the OE of \textit{Xenopus laevis} tadpoles and mammals

In this thesis, it was demonstrated that the action of 2-AG controls detection thresholds of odorants in larval \textit{Xenopus laevis}. Subsequently, it was investigated, whether the OE of higher vertebrates also exhibit the equipment of the endocannabinoid system. For this experiment, samples of rats and a human were used.

The mRNA for the enzymes for production and degradation of 2-AG as well as anandamide were detected in rat and human. The CB1 and CB2 receptor were only found in rats, but GPR55 was detected in both species (Table 6). GPR55 is an orphan G protein-coupled receptor and was recently proposed as a novel cannabinoid receptor with different pharmacological properties than CB1 and CB2 receptors (Barnett-Norris \textit{et al.}, 2005; Brown, 2007; Pertwee, 2007). Hence, the components or similar components of the endocannabinoid system found in larval \textit{Xenopus laevis} also exist in the OE of rat and human. In order to speculate about similarities between the physiological meaning of endocannabinoids in the OE of larval \textit{Xenopus laevis} and mammals, further experiments have to be done.
Recently, several studies revealed other neuromodulatory substances like orexin, leptin, adiponectin, and dopamine. Orexin, leptin, and adiponectin are hormones involved in food intake and energy metabolism, and act orexigenic or anorexigenic. Orexin and leptin are synthesized in the OE (Caillol et al., 2003; Baly et al., 2007) whereas adiponectin reaches the OE with the blood serum (Hass et al., 2008). Dopamine acts neuroprotectively in the OE, probably by binding to D2 receptors (Hegg and Lucero, 2004). The source of dopamine is not identified yet. However, all described substances have their corresponding receptors on ORNs and probably influence olfactory transduction. Odorant transduction and coding seems to be a complex, highly dynamic process. Olfactory receptor neurons are thus not only tuned by 2-AG, as demonstrated in detail in this thesis, but also by many other substances. To gain first hints if these systems indeed exist in parallel within the OE of one animal, mRNA of OEs of larval Xenopus laevis, rats, and the human were screened for receptors of orexin, leptin, adiponectin, and dopamine.

In tadpoles, mRNA for dopamine, adiponectin, and leptin receptors were found (the genetic identity for the orexin receptor is not known so far). Rats exhibit mRNA for dopamine, adiponectin, orexin, and leptin receptors. The human sample showed adiponectin as well as leptin receptor mRNA, but neither dopamine nor orexin receptor mRNA (Table 6). These data demonstrate that the morphological basis for a complex modulatory signaling system is present in all three species. To date it is not known how these systems act, and whether they interact with the endocannabinoid system. However, in the literature there are some hints for possible interplays of these systems:

Peptides such as leptin, adiponectin, and orexin modulate food intake, and act on the endocannabinoid system. Consistently, it was shown that leptin treatment of mice affects 2-AG signaling in the thalamus (Di Marzo et al., 2001; Jo et al., 2005). A functional link between adiponectin and endocannabinoids was observed by Zyromski et al. (2009). In that study, CB1 receptor blockade increased circulating adiponectin concentration. Crespo and coworkers (2008) demonstrated that the CB1 receptor antagonist rimonabant blocks the orexigenic effect of orexin. The receptor for orexin belongs to the family of G_{q/11} protein-coupled receptors (Sakurai et al., 1998). These receptors have the potential to stimulate endocannabinoid production through receptor-driven endocannabinoid release or Ca^{2+}-assisted
receptor-driven endocannabinoid release (Hashimotodani et al., 2007; Kano et al., 2009). Thus, orexin may be a trigger for endocannabinoid release in the OE.

Another type of interaction between dopamine and orexin on the one hand and endocannabinoids on the other is receptor heterodimerization of the respective receptor with the CB1 receptor (Mackie, 2005). CB1 receptors and D2 receptors can form dimers with antagonistic interactions (Marcellino et al., 2008). Furthermore, CB1 and orexin receptors can dimerize (Hilairet et al., 2003; Ellis et al., 2006).

Many modulatory systems exist in the OE. Their roles and their possible interfaces have to be investigated by physiological experiments in future studies. The presence of various modulatory systems in the OE and the few studies indicating first functional links open the novel field of peripheral modulation of olfactory input.

4.2.6 Conclusions

To summarize, the findings support the view that paracrine 2-AG acts as an orexigenic modulator in the OE by increasing and decreasing the sensitivity of ORNs to odorants during phases of hunger or satiety (Figure 26). As a consequence, the concept of a “threshold as a well defined and constant concentration below which an ORN does not respond to a specific odorant” has lost its meaning. PMCA probably acts as the effector molecule of the CB1 receptor and alters the intracellular Ca$^{2+}$ concentration according to the 2-AG level in the OE. This would be a novel signaling mechanism of the CB1 receptor and a future study will reveal the cascade in detail. Besides endocannabinoids, also other modulatory substances may act in the OE.
Figure 26: Scheme of endocannabinoid action in the OE. Under control conditions a tonic level of 2-AG is synthesized. DAGLα mRNA synthesis in sustentacular cells (~, blue) is enhanced upon food-deprivation and leads to an enhanced level of 2-AG binding to CB1 receptors (green circles) on ORN dendrites (blue arrows). This state renders ORNs more sensitive and increases action potential frequency. DAGLβ mRNA expression in ORNs (~, red) is not affected upon food-deprivation. 2-AG synthesized in ORNs feeds back on ORNs (red arrow). The CB1 receptor probably alters PMCA activity. Also other substances like orexin, leptin, adiponectin, and dopamine may modulate ORN activity. PC, principal cavity; ON, olfactory nerve; od, odorant; SC, sustentacular cell.
5 Summary

The sense of smell is critical for finding food and mediating emotional and social responses. A dysfunction of this sense is often associated with diseases. In the last two decades much progress has been made in this research field and insights in how the sensation of smell is perceived has been gained. This thesis covers two topics of basic research in olfaction and is accordingly subdivided:

1. Various olfactory receptor neuron subsets exist in the olfactory epithelium. One classification of these subsets can be made according to the transduction mechanism of odorants. The olfactory receptor neurons belonging to one subset transduce odorants into depolarizations using the cAMP-dependent transduction mechanism. To date, it is not possible to stain this subset or to interfere with their generator channels with potent and specific blockers. In this thesis, the styryl dye FM1-43 was identified as a marker for olfactory receptor neurons endowed with cAMP-dependent transduction machinery. The dye is internalized in neurons by uptake through cyclic nucleotide-gated channels, the generator channel of the cAMP-dependent olfactory receptor neurons. This was proven by interference of FM1-43 uptake with divalent ions and unspecific cyclic nucleotide-gated channel blockers used in high concentrations and further confirmed by evoking responses to cAMP and forskolin in FM1-43-stained cells. Characteristic for FM1-43-stained olfactory receptor neurons is that these cells did not respond to odors or rapidly lost their responsiveness. This suggested that FM1-43 severely interfered with the transduction machinery of stained olfactory receptor neurons. Indeed, 10 µM FM1-43 blocked currents through native cyclic nucleotide-gated channels to approximately 25 % and acts from the extracellular side. This tool thus allows optical differentiation and pharmacological interference with olfactory receptor neurons endowed with cAMP-dependent transduction at the level of the signal transduction.
Summary

2. The sense of smell is an important input for the search for food and food intake, but the underlying mechanisms for this functional interaction are poorly understood. One key factor for energy homeostasis and nutrition at central stages is the endocannabinoid system. Therefore, it was hypothesized, that the endocannabinoid system may link food intake with olfaction. Recently, cannabinoids were shown to act in the olfactory epithelium. Pharmacological interference at the according cannabinoid receptor modulates odor-evoked responses. In this thesis, it is shown that the endocannabinoid 2-AG is synthesized in the olfactory epithelium and acts on olfactory receptor neurons. Blocking 2-AG synthesis decreased and delayed odorant-induced responses. Analyzing single cells revealed, that there are two sources of 2-AG in the olfactory epithelium: the first are olfactory receptor neurons, where production of 2-AG depends on diacylglycerol lipase $\beta$, and the other are sustentacular cells, where production depends on diacylglycerol lipase $\alpha$. Diacylglycerol lipase $\alpha$ mediated 2-AG-synthesis in sustentacular cells is influenced by the hunger state of the animal. The essential 2-AG effect in olfactory receptor neurons is the control of odorant detection thresholds. An enhanced 2-AG level decreases the detection threshold of an individual olfactory receptor neuron whereas a lowered 2-AG level increases the detection threshold. Thus, hunger renders olfactory receptor neurons more sensitive and endocannabinoid modulation in the nose may therefore substantially influence food seeking behavior. The intracellular effector mediating cannabinoid receptor actions probably is the plasma membrane calcium ATPase. This signaling cascade was not described as yet. However, future experiments have to be performed to identify the complete transduction cascade. Besides the endocannabinoid system, receptors for several other modulatory substances, i.e. orexin, leptin, adiponectin, and dopamine, were found in the olfactory epithelium. This indicates that already at the most peripheral stage of the olfactory system, the sensation of odors is modulated by many substances.
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