Differential Effects of Acetylcholine on Neuronal Activity and Interactions in the Auditory Cortex of the Guinea-pig

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Abstract

During normal brain operations, cortical neurons are subjected to continuous cholinergic modulations. In vitro studies have indicated that, in addition to affecting general cellular excitability, acetylcholine also modulates synaptic transmission. Whether these cholinergic mechanisms lead to a modulation of functional connectivity in vivo is not yet known. Herein, the effects were studied of an iontophoretic application of acetylcholine and of the muscarinic agonist, carbachol, on the ongoing activity and co-activity of neurons simultaneously recorded in the auditory cortex of the anaesthetized guinea-pig. Iontophoresis of cholinergic agonists mainly affected the spontaneous firing rates of auditory neurons, affected autocorrelations less (in most cases their central peak areas were reduced), and rarely affected cross-correlations. These findings are consistent with cholinergic agonists primarily affecting the excitability of cortical neurons rather than the strength of cortical connections. However, when changes of cross-correlations occurred, they were usually not correlated with concomitant changes in average firing rates nor with changes in autocorrelations, which suggests a secondary cholinergic effect on specific cortico-cortical or thalamo-cortical connections.

Introduction

The neocortex is innervated by cholinergic fibres that originate in the basal forebrain (reviewed in Mesulam, 1995). In the auditory cortex of the rat (Lysakowski et al., 1989), cat (Irlé and Markowitsch, 1984) and monkey (Mesulam et al., 1983), the presence of cholinergic innervation was demonstrated by using different cholinergic markers, such as acetylcholinesterase (AChE, the degradative enzyme of acetylcholine (ACh)) and choline acetyltransferase (ChAT, the synthesizing enzyme of ACh). The majority of the cholinergic projections to the cortex originate in extracortical structures, mainly in the nucleus basalis magnocellularis (Mesulam, 1995).

The possible implication of ACh in cortical sensory processing has become of considerable interest because of ACh's involvement in behavioural processes related to selective attention, arousal, motivation and reward (reviewed in Semba, 1991), as well as in cellular plasticity (Hars et al., 1993; Auerbach and Segal, 1994; Ahissar et al., in press). The role of ACh in these processes has been analysed at the biophysical and functional levels in different species and cortical sensory areas (McCormick, 1992a). The iontophoretic application of ACh or the stimulation of the basal forebrain result in a muscarinic-mediated modulation of firing modes (McCormick and Prince, 1986; Metherate et al., 1992; Cox et al., 1994). Studies of the modulation by ACh of receptive field properties of sensory cortical neurons for different modalities (Silitto and Kemp, 1983; Sato et al., 1987; Lamour et al., 1988; Metherate et al., 1988; Bassant et al., 1990) generally show a facilitatory influence of muscarinic agonists on cortical neurons, with varying proportions of cells inhibited by ACh.

This latter inhibitory effect is usually attributed to the excitation of intracortical inhibitory interneurons (McCormick and Prince, 1986; Muller and Singer, 1989). Despite a wealth of studies focusing on the effects of ACh on ongoing activity and sensory responses at the level of the single cell, studies on the possible modulatory effects of ACh at the level of intercellular communication are lacking. It is usually assumed that most action potentials discharged by cortical neurons are the result of a joint effect of correlated inputs (Abeles, 1982b). Modulation of correlated activity could be achieved by activation of the cortical cholinergic basal forebrain system, which is known to change the global activity state of the cortex (Semba, 1991) and the thalamo-cortical transmission gain (Metherate and Ashe, 1993). The various effects of ACh that have been described at the biophysical level, usually in in vitro preparations, predict different outcomes on the network activity and cross-interactions between cells. While the modulation of excitatory synaptic transmission (Markram and Segal, 1990, 1992) predicts increased levels of functional coupling between neurons, which result in turn in changes in firing activity, the effects of ACh on excitability and firing modes (McCormick and Prince, 1986) predict a different scenario in which the firing rates will be primarily affected by ACh in vivo and interneuronal interactions will be affected to a lesser degree, if at all.

In this study, we compared the effect of ACh on neuronal excitability and functional connectivity. The effects of an iontophoretic application of ACh or of a muscarinic agonist, carbachol, on the ongoing firing...
rates and auto- and cross-correlations were studied in the auditory cortex of the anaesthetized guinea-pig. Preliminary accounts of these data have been presented in abstract form (Cohen et al., 1994; Shulz et al., 1995).

Materials and methods

Histochemical localization of acetylcholinesterase

A standard histochemical staining procedure (Biegon et al., 1986) was used to localize AChE in the auditory cortex. Briefly, an adult guinea-pig was decapitated, and its brain was quickly removed, blocked in the plane of section, and frozen on dry ice. Two series of interleaved coronal sections of 30 mm were made with a cryostat and collected on gelatin-coated slides. One series was used for Nissl staining and the other for AChE histochemistry. For the latter, the sections were fixed in PBS and incubated (2 h, 37°C) in a 0.05 M acetate buffer (pH 5) containing 10 mM glycine, 2 mM copper sulphate, 4 mM acetylthiocholine iodide, and 10 mM ethopropazine. The stain was developed for 1 min at room temperature in 1.25% sodium sulphide at pH 7.5 and intensified for 1 min in a 1% silver nitrate solution. Finally, the slides were transferred to phosphate-buffered formalin for 30 min, dehydrated through a series of alcohols, cleared in xylene, and coverslipped. Distribution of the AChE staining was determined by microscopic examination and compared to the cytoarchitectonic structure of the cortex revealed with the Nissl staining.

Preparation

Thirty-eight Dunkin–Hartley guinea-pigs (Cavia porcellus Linnaeus) weighing 380–940 g (median, 580 g) were used in these experiments. The animals were anaesthetized by an i.p. injection of urethane (1.2 g/kg) followed by an i.m. injection of Combelen (propionylpropamine hydrochloride, 3.6 g/kg). Surgical levels of anaesthesia were then achieved by an i.m. injection of Rompun (xylazine hydrochloride, 8 mg/kg). Throughout each experiment, supplementary i.m. injections of Ketalar (ketamine hydrochloride, 100–200 mg/kg) were administered to maintain an adequate level of anaesthesia. In a typical experiment, recording from cortical units was not started for at least 2 h after the initial induction of anaesthesia.

The animal was mounted on a head-holder without ear bars, which permits free access to the auditory cortex and avoids damage to the tympanum membrane (Haidarliu, 1996). A local anaesthetic (lidocaine, 2%) was applied to all wounds. A small hole (3 mm diameter) was made in the skull to expose the auditory cortex, and the dura was removed. Drying of the surface of the cortex was prevented by repeated topical applications of saline. The animal was placed in a sound-attenuating chamber for the recording session.

Electrodes and drug applications

The multi-electrode setup used for combined micro-iontophoresis and multiple single-unit recordings has been described previously (Haidarliu et al., 1995). Briefly, the setup consists of two types of electrodes included within a metallic guide tube: regular tungsten-in-a combined electrode (CE), which consisted of a tungsten electrode (0.2–0.8 MΩ) within the central barrel of a seven-barrelled glass pipette. Within the stainless steel guide tube, the regular and combined electrodes were assembled in two different configurations: (i) one CE and three TEs, referred to as the ‘single combined’ configuration (SC), and (ii) two CEs and two TEs, called the ‘double combined’ configuration (DC). Depending on the configuration, the horizontal distance between the electrode tips was 300–600 μm. The guide tube was brought close to the exposed brain, and each of the four electrodes was introduced independently into the auditory cortex with a multi-electrode microdrive system. Four spike sorters (MSD-2; Alpha-Omega, Nazareth, Israel), one for each electrode, were used to isolate single- and multi-unit spikes. The separations of units from a given electrode were classified according to the following five categories: well isolated units (q5), well isolated units of which up to 10% of the spikes were not detected (q4), two-single units that could not be separated reliably on the basis of their templates (q3), multi-units (q2), and multi-units with amplitudes too close to noise level (q1; these were excluded from the analysis). Hereafter, ‘well-isolated units’ refers to q4 or q5, ‘isolated units’ to q3, q4 and q5, and ‘multi-unit’ to q2 units. The templates and the recorded action potentials were carefully and continuously inspected with both the spike-sorter software and the oscilloscope. Recording of isolated single units was usually stable for several hours (Fig. 4 in Haidarliu et al., 1995). Whenever a deviation from the steady state was noticed, measures to correct the deviation were taken, including changes in the sorting threshold and template updating. In extreme cases, the electrode was moved a few micrometres to restore the original recording situation. The data collected during these periods of template recovery were excluded from analysis. If restoration of the original template could not be achieved, the cell recording was stopped.

For the iontophoretic application of neuroactive drugs, the CE was filled with the different solutions before insertion into the guide tube. The six barrels were filled with an aqueous solution of ACh chloride (1 M, pH 4.5), carbamylcholine chloride (carbachol; 1 M, pH 6.0), atropine sulphate (0.1 M, pH 4.5), (—)artcarnel bitartrate (noradrenaline; 0.5 M in 100 mM ascorbic acid solution, pH 4.5), or NaCl (3 M). All these solutions were frozen and stored at —20°C as aliquots and were thawed immediately before the experiment. The impedance of the glass barrels varied from 10 to 100 MΩ (median, 30 MΩ), depending on the diameter of the capillary orifice and on the filling solution. Retaining currents of —5 nA were used to prevent the drugs from leaking. The ejection currents used were 20—200 nA (positive current). Balanced ejections were not systematically applied in order to avoid ejection of the active drug by the balance barrel during retention periods (Sillito and Kemp, 1983). No qualitative difference was found between balanced and unbalanced applications. The use of high current levels to affect the activity of units recorded far from the site of ejection increases the possibility that some of the effects observed might result from current effects. However, most of the drug applications were in the 20—100 nA range, and the time course of the effects was generally too slow to be caused by direct current effects. Moreover, the blockade of the effects by atropine (see Results) strengthens the probability of a specific effect of cholinergic agonists via muscarinic receptors.

Data analysis

The modulation of the firing rate by ACh or carbachol was assessed as follows. The firing rate was plotted as a function of time (a ‘rate meter’) with a resolution (bin size) of 20 s. This function was visually inspected for deviations from baseline that occurred during or immediately after the drug application. A deviation was considered significant if it was unmistakably larger and/or consistently longer than any other deviation prior to the application. In cases where a given unit received more than one drug application, the unit was considered cholinceptive only if the effect (deviation) was reproducible. A statistical value was assigned to these visually judged effects by the following calculation (see also Metherate et al., 1988). First,
120 recorded units were selected from the total of 235 recorded units in which the spontaneous period before application was long enough (at least 300 s) to estimate the relationship between the average and the standard deviation (SD) of the firing rate (for the calculation of the SD, this spontaneous period was divided in 6 bins of 50 s each). A single regression function could be fitted to the data \[ \log(\text{SD}(R)) = (0.69 \pm 0.04) \times \log(\text{Avg}(R)) - (1.46 \pm 0.05), \] where \( n = 767, r = 0.74, P < 0.001 \). Then, for each of the 235 cells and for each application of ACh, the expected SD was estimated according to the regression function and the average firing rate of the cell during the spontaneous period preceding the application. Any modulation of the firing rate, during or soon after the application of the drug, was considered as significant if the firing rate differed from the expected firing rate by 2 or more SDs. For 80% of the visually selected significant effects, the deviations were larger than \( \pm 4 \) SD of the mean firing rate.

Auto- and cross-correlation histograms (Perkel et al., 1967a, b; time bins of 1, 2 or 5 ms) were calculated for spike trains recorded during periods of spontaneous activity. The correlograms were tested by (i) visually inspecting for qualitative changes in shape, and (ii) quantitatively analysing the central peaks of the histograms within two time-lag ranges (0–20 and 0–300 ms). For each histogram, the measure, known as the asynchronous gain (asg; Levick et al., 1972; Abeles, 1982b; Vaadia et al., 1991), was used to estimate the functional coupling between neurons. Two ‘asg meters’, which describe the evolution of the asg value as a function of time in 50 s intervals, were computed for each pair of simultaneously recorded neurons. The modulations of the asg-meters due to ACh or carbachol applications were studied and tested statistically with the same method that was used for rate meters. The corresponding regression functions for auto- and cross-correlograms were:

\[ \log(\text{SD}(\text{asg}_\text{auto})) = (0.38 \pm 0.03) \times \log(\text{Avg}(\text{asg}_\text{auto})) \times (1.06 \pm 0.05), \] where \( n = 767, r = 0.74, P < 0.001 \) and

\[ \log(\text{SD}(\text{asg}_\text{cross})) = (0.43 \pm 0.02) \times \log(\text{Avg}(\text{asg}_\text{cross})) - (0.69 \pm 0.05), \] where \( n = 767, r = 0.66, P < 0.001 \).

The dependency of the results on the specific normalization of the peak area was tested by repeating all the calculations with the modified correlation coefficient \( \text{cc}' \), which equals the area under the peaks of the two units (Lavner, 1989). Although for any given neuronal pair the \( \text{cc}' \) values differed from the \( \text{asg} \) values, significant differences for the population as a whole were not found when measuring the effects of cholinergic applications.

### Results

#### Staining for AChE in the auditory cortex

Specific staining for AChE was evident in all neocortical structures of the guinea-pig, as well as in the thalamus; the strongest staining was observed in the hippocampal formation (Fig. 1A). The distribution of AChE staining across layers in the cortex was determined by microscopic examination of matching AChE- and Nissl-stained sections. The layering pattern of staining appeared to depend on the cortical area. For example, the perirhinal area, which is ventral to the auditory cortex in Figure 1, exhibited stronger staining in the supragranular layers than in the auditory cortex. The organization of the AChE labelling of the auditory cortex is illustrated in Figure 1. Two prominent dark bands of AChE staining are manifested; the stronger one is confined to layer I and the second occurs at infragranular layers V and VI. Layer IV was more lightly stained than other layers in the cortex. The background staining in the white matter was very light, with apparently AChE-positive fibres passing through it. In these coronal sections, only short segments of the axons were...
visualized, which suggests that AChE positive axons run parallel to
the antero-posterior axis in cortical white matter.

Cholinergic electrophysiological effects

Two hundred and ninety one units were recorded in the auditory
cortex of 38 guinea-pigs. The response to the iontophoretic application
of ACh or carbachol was tested for all units. However, those
penetrations in which none of the recorded cells responded to the
applied drugs, even at high levels of current, were excluded from the
statistical analyses. A total of 56 cells were excluded (nine experi-
ments, ten penetrations), and the database was restricted to 235 units,
of which 150 were recorded with the SC electrode configuration and
85 cells with the DC configuration. The exclusion probably led to an
overestimation of the proportion of cholinceptive cells, but was used
since there was no indication in the excluded cases that the drug was
effectively ejected from the pipette. In all these experiments, only
one penetration (with the four electrodes simultaneously) was made
per animal, and lasted on average 10 h; in only three cases were two
different groups of units successively recorded during a given
penetration. According to the quality of the discrimination of action
potentials from a given electrode, 94 recordings were from well
isolated units (q4 and q5), 72 from two single units (which could not
be separated reliably on the basis of their templates, q3), and 69 from
multi-units (q2). Statistical analyses were performed on both the total
and restricted samples, as a function of electrode configuration,
distance from the site of ejection, and spike separation quality (see
below). In addition to ACh, the agonist carbachol was iontophoresed,
since its slower hydrolysis rate increases the probability of diffusion
over larger volumes of cortical tissue. The percentage of units that
exhibited significant modification of their spontaneous activity upon
separate exposure to both drugs is shown in Table 1. Cells were not
tested with both ACh and carbachol. However, since the effects of
the two agonists did not differ qualitatively or quantitatively in the
percentage of modified cells ($\chi^2, P = 0.134$), the data obtained
separately with both agonists were pooled for subsequent analysis.

Effects of cholinergic agonists on the spontaneous activity
rates

The cholinergic agonists affected the ongoing activity of 40.5% (95/
235) of the cells tested. In 25.5% of all the cases the spontaneous
activity was significantly increased, whereas in 15% of the cases it
was reduced. The mean current levels used for inducing excitatory
and inhibitory effects were not significantly different. In a few cases
($n = 7$), the drugs induced biphasic responses (Fig. 2).

A clear relationship was not observed between the initial level of
spontaneous activity and the probability of affecting the ongoing
activity. Quiescent cells, which had 0—0.5 spikes/s, did not appear
to be less readily excited by ACh or carbachol (19/81, 23.5%) than
cells with spontaneous activity of 0.5—20 spikes/s (40/154, 25.9%),
as previously reported in the somatosensory cortex (Bassant et al.,
1990). The probability of a rate decrement showed an initial trend;
cells with lower initial levels of spontaneous activity were less
affected than cells with higher initial activity. This trend, which could
be due to the lower detectability of decrements for initial low firing
rates, holds for discharge frequencies up to five spikes per second
and reverts above it. When the analysis was restricted to isolated
units, the same relationship between the spontaneous rate and the
probability of being affected by ACh application was observed both
for rate increments and decrements. Multunit recordings showed a
smaller, but still not significantly different, percentage of modification
than well-isolated units (24.5% versus 39.6%, $\chi^2, P = 0.1$; Table 2).

Effect on neuronal activity of the distance between the
recording and iontophoresis sites

Depending on the electrode configuration (SC or DC), one or two
sites of ejection were used, respectively. For each configuration, two
groups of cells were defined, those recorded from the electrode
through which the drugs were applied (near cells) and those recorded
by one of the other electrodes at a variable distance (usually 300—
600 $\mu$m) from the site of application (far cells). Herein, the extent of
the modulatory action between the two groups of cells is compared
without considering the exact distance between the tips of the
electrodes.

The population of near cells recorded with the SC configuration is
the one that should be compared with cells in previous studies in
which one electrode was used for both recording and ejection. The
percentage of isolated (single or two) units with a modification of
the ongoing firing rate was determined as a function of the recording
site relative to the application site when the SC setup was used (Table
3). Iontophoresis of cholinergic agonists affected (i.e. caused rate
increments, decrements or biphasic effects) almost twice as many
isolated units ($n = 101$) recorded with the CE (61.8% of near cells,
$n = 21$) than cells recorded simultaneously through one of the other
electrodes (33.7% of far cells, $n = 80$, $\chi^2, P < 0.036$). The proportion of cells that exhibited a facilitatory effect was similar for
far and near groups (17.5 and 19% respectively of the total units).
The proportion of far cells (13.7%) that exhibited a decreased level
of spontaneous activity was less than that of near cells (33.3%). This
is consistent with an ACh-induced decrease in spontaneous activity
being mediated by the activation of local GABAergic interneurons
(McCormick and Prince, 1986; Muller and Singer, 1989). An interneu-
ron effect on far cells is expected to be less effective, since (i) local
interneurons do not directly drive far (300—600 $\mu$m) neurons, and
(ii) far interneurons are probably only weakly affected by the diffused
cholinergic agent, and this effect is insufficient to be expressed in
the second-order neuron’s activity.

Involvement of muscarinic receptors

The receptors involved in mediation of the cortical effects of
cholinergic agonists were examined by simultaneously ejecting a
cholinergic agonist and antagonist through the same and/or different
electrodes of a DC setup. The firing rate of nine simultaneously
recorded units as a function of time is shown in Figure 2; five of
these units were recorded by the CE electrodes (units 1, 2, 3, and 8,
9) and four by the TE electrodes (units 4, 5, 6 and 7). Carbachol was
repeatedly applied through one of the two CEs (electrode 1), and
excited several units, maximal significant effects being observed with
those units recorded near the site of ejection (units 1, 2 and 3). The
responses of units far from the site of ejection (e.g. units 4, 5 and 8)
were usually weaker, delayed, and less consistent, i.e. the response
pattern did not repeat for every application, and then was not
considered as significant. Units 6 and 7 showed spontaneous modifica-

<table>
<thead>
<tr>
<th>Changes in firing rate</th>
<th>Application of Acetylcholine</th>
<th>Carbachol</th>
<th>Acetylcholine or Carbachol</th>
</tr>
</thead>
<tbody>
<tr>
<td>Increased</td>
<td>16 (18%)</td>
<td>44 (30%)</td>
<td>60 (25.5%)</td>
</tr>
<tr>
<td>Decreased</td>
<td>14 (16%)</td>
<td>21 (14%)</td>
<td>35 (15%)</td>
</tr>
<tr>
<td>Unchanged</td>
<td>58 (66%)</td>
<td>82 (56%)</td>
<td>140 (59.5%)</td>
</tr>
<tr>
<td>Total number of cells</td>
<td>88</td>
<td>147</td>
<td>235</td>
</tr>
</tbody>
</table>

TABLE 1. Modulation of firing rates upon exposure to acetylcholine or carbachol


tions of firing rates just preceding the first application. Neither unit was considered as showing a significant effect. Unit 9 did not show any significant effect. Application of atropine through the second CE (electrode 4), which was several hundred micrometres from the site of ejection of carbachol, partially and reversibly blocked the effect, thus suggesting that atropine diffused several hundreds of microns to block the muscarinic receptors on the membrane of the cells recorded by the other CE (electrode 1). When atropine was applied simultaneously with carbachol through the same electrode, the effect of carbachol was completely blocked and the blockade lasted ~50 min (see last application of carbachol). Since the effects of iontophoresis of ACh or carbachol could be completely blocked by a simultaneous application of atropine, the cortical effects of these cholinergic agonists appeared to be mediated through muscarinic receptors.

Cholinergic effects on firing patterns assessed by autocorrelations

At short time lags (a few to a few tens of milliseconds), the shape of an autocorrelation histogram usually reflects the tendency of the cell to fire in regular or bursting modes. Accordingly, the autocorrelation histograms were quantified for 235 cells before, during and after the application of ACh or carbachol by calculating the asg values for a 20 ms time lag. Since many of the neurons recorded in this study usually exhibited a peak of a few hundreds of milliseconds in the autocorrelation, the asg values for a 300 ms time lag were also calculated.

The percentage of autocorrelation histograms significantly affected by the application of ACh or carbachol was 9.4% (22/235) and 10.6% (25/235) for the 20 and 300 ms time lags, respectively. Only eight of the units having their autocorrelation significantly affected for the 20 ms time lag were also significantly affected for the 300 ms time lag. By restricting the analysis to isolated units whose initial spontaneous activity was at least 0.5 spikes/s (below this firing rate the number of action potentials was not sufficient for estimation of the autocorrelation function), the percentage of affected cells increased to 13.6% (15/110) and 16.4% (18/110), respectively. In the majority of cases where a modification was observed, a significant decrease in the probability of firing within both delay periods was observed [60% (9/15) for the 20 ms time lag and 77.8% (14/18) for the 300 ms time lag]. The autocorrelograms of five representative cells, four of which showed a significant decrease in the area of the histogram at 0—20 ms, are depicted in Figure 3. One cell (Fig. 3E) displayed an increase in its tendency to fire bursts of spikes upon application of ACh. For this latter cell, the absolute refractory period (4 ms) was unchanged, while the relative refractory period was shorter and the probability of firing after each action potential during the first 20 ms increased. ACh decreased the probability of repetitive firing by 0.29 ± 0.17 (SD) spikes on the average of the cells that exhibited a reduction of the asg (n = 13) in the 0—20 ms delay. The cells that showed an increase of the asg (n = 9), upon application of ACh tended to fire an extra 0.32 ± 0.18 spikes during the first 20 ms.

In the majority of cases (93 and 83% for the 20 and 300 ms time lag groups, respectively), changes in the autocorrelograms were concomitant with changes in the firing rate levels. The occurrence of changes in the autocorrelograms was significantly correlated with the occurrence of rate changes (χ², P < 0.001) and P < 0.007 for the 20 and 300 ms time lag groups, respectively. However, the directions of the changes were not correlated (χ², P = 0.69 and P = 0.78 for the 20 and 300 ms time lag groups, respectively).

Cholinergic effects on functional coupling assessed by cross-correlograms

Cross-correlations measure the net effect, mediated by synaptic connections, of one cell on another. Thus, changes in cross-correlations reflect changes in the underlying synaptic weights, and steady cross-correlations reflect either steady synaptic weights or balanced changes within the synaptic array.

For the 235 units analysed with either of the two electrode setups, 1710 cross-correlations were computed, as were the asg values for two regions of time lags (20 and 300 ms) during periods of spontaneous activity and during the application of ACh or carbachol. These time lags were chosen since similar time lags were used for previously reported distributions of cross-correlogram peak widths (Krüger and Aiple, 1988; Lavner, 1989; Nelson et al., 1992). The number of pairs of cells whose functional coupling, as assessed by asg values, changed significantly upon application of cholinergic agents was extremely low; only 3.2% (54/1710) and 4.3% (73/1710) of the cross-correlograms were affected within the 20 and 300 ms time lags, respectively. These percentages remained low (3.3 and 5.6% respectively, n = 642) even if only isolated units with spontaneous activity levels such that the product of the individual firing rate was higher than 0.25 spikes/s were considered. Furthermore, significant differences were not observed for pairs in which at least one cell was recorded near the site of iontophoresis or in which both cells were recorded by the far electrodes. The probability of modification was similar for the DC and SC setups. With the DC configuration, the distance of recording, the quality of the unit isolation, and even the initial firing rate did not significantly affect the probability of inducing a change in the functional coupling. Of the 54 pairs of cells whose asg values for the 20 ms delay changed, 43 (79.6%) had an increase in the asg value. Only ten pairs of units showed significant modulations of the asg values for both 20 and 300 ms time lags.

An example of a one-sided peak that appeared immediately after the application of ACh is depicted in Figure 4. The units of the pair were recorded by two different electrodes, and the supposedly postsynaptic unit (cell 11) was recorded near the site of ejection. The asg was increased by ACh by >8 SD (peak asg, 0.11; spontaneous asg, 0.007 ± 0.012). The significant change in the cross-correlogram occurred in the absence of a noticeable change in the firing rate or pattern of either unit, as shown by the unchanged autocorrelograms and rate meters in Figure 4.

Whether the detection of a change in the cross-correlogram depends on the spontaneous level of correlation was tested by computing the percentages of modified pairs as a function of their initial spontaneous correlation level (Table 4). To avoid artificial bias due to few
Cholinergic modulation of correlated activity

ATROPINE
CARBACHOL (e1)

ELECTRODE 1
unit 1, q2
unit 2, q3
unit 3, q5

ELECTRODE 2
unit 4, q2
unit 5, q2

ELECTRODE 3
unit 6, q2
unit 7, q4

ELECTRODE 4
unit 8, q2
unit 9, q3

Graph showing units 1, 2, and 3 with time in seconds and spikes per second, and a scale for 0.5 ms and 150 μV.
402 Cholinergic modulation of correlated activity

Table 2. Effects of acetylcholine and carbachol on firing rate as a function of spike discrimination

<table>
<thead>
<tr>
<th>Firing rate</th>
<th>Spike discrimination</th>
<th>All</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Isolated units (q3, q4, q5)</td>
<td>Multi-units (q2)</td>
</tr>
<tr>
<td>Increased</td>
<td>18 (17.8%)</td>
<td>6 (12.2%)</td>
</tr>
<tr>
<td>Decreased</td>
<td>18 (17.8%)</td>
<td>3 (6.1%)</td>
</tr>
<tr>
<td>Biphasic</td>
<td>4 (4.0%)</td>
<td>3 (6.1%)</td>
</tr>
<tr>
<td>Unchanged</td>
<td>6 (60.4%)</td>
<td>37 (75.5%)</td>
</tr>
<tr>
<td>Total number of units</td>
<td>101</td>
<td>49</td>
</tr>
</tbody>
</table>

SC configuration used to record all units.

Table 3. Effects of acetylcholine and carbachol on the firing rate of isolated units as a function of proximity to the application site

<table>
<thead>
<tr>
<th>Firing rate</th>
<th>Application site</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Near</td>
</tr>
<tr>
<td>Increased</td>
<td>14 (17.5%)</td>
</tr>
<tr>
<td>Decreased</td>
<td>11 (13.7%)</td>
</tr>
<tr>
<td>Biphasic</td>
<td>2 (2.5%)</td>
</tr>
<tr>
<td>Unchanged</td>
<td>53 (66.3%)</td>
</tr>
<tr>
<td>Total number of units</td>
<td>21</td>
</tr>
</tbody>
</table>

SC configuration used to record all units.

outliners, this analysis was restricted to the population of pairs whose spontaneous absolute asg values were <3 SDs of the total population. The total population had means and SDs of 0.034 ± 0.103 for the 20 ms and 0.162 ± 0.238 for the 300 ms time lag. The restricted population included pairs whose lasgl < 0.3 (asg = 0.022 ± 0.045) for the 20 ms time lag and lasgl < 0.7 (asg = 0.11 ± 0.179) for the 300 ms time lag. The SDs of the spontaneous asg values of these restricted populations were used to categorize the spontaneous asg levels of single pairs. The percentage of affected pairs increased as the spontaneous initial correlation increased, up to ~2 SDs, and then decreased. It is worth noting, however, that even the highest percentages of affected pairs (6.7% for the 20 ms and 8.1% for the 300 ms time lag) were still significantly lower than the percentage of units whose firing rates were affected. In only ~1% of the pairs that showed small spontaneous correlation (<0.5 SD; Table 4) did cholinergic applications induce a significant change (Fig. 4).

An example of a clear change in cross-correlation for a pair of units with an initial spontaneous correlation is depicted in Figure 5. Three successive cholinergic applications induced three successive increments in the peak area of the cross-correlation between the two neurons, which were recorded by different electrodes. These changes were accompanied by concomitant increments of the peak area of the autocorrelation of the target neuron (unit 2) and decrements in the background level (flanks) of the autocorrelation of the trigger unit (unit 4).

No systematic correlation was observed between changes in the asg values of the cross-correlograms and changes in the firing rates and patterns (autocorrelograms). The distribution of modified auto- and cross-correlations is shown as a function of concomitant changes in the firing rates and patterns (Fig. 6). While most of the changes in the autocorrelograms were accompanied by changes in the levels of spontaneous activity, the latter was not a reliable predictor of a change in the cross-correlation. For example, 33% of the changes in the cross-correlograms occurred without any change in firing rate (Fig. 6C, D, left columns). Moreover, two thirds of the changes in the cross-correlograms occurred without any significant change in either of the autocorrelograms (Fig. 6C, D, middle columns), while a third of the changes in cross-correlograms occurred without any change in the firing rate and in the autocorrelograms of both units of the pair (Fig. 6C, D, right columns).

The relative effect of ACh on firing rate and auto- and cross-correlations was assessed by computing the percentages of affected units or pairs of units using different criteria. The cases were grouped according to which electrode setup (SC or DC) and site of recording (near or far) was used. For cross-correlations, cases in which both neurons were recorded far from the site of iontophoresis were considered as ‘far,’ and all other cases as ‘near’. These groups were further reduced to sets that included only isolated units (g > 3), and then to sets containing only isolated units with a spontaneous firing level ≥0.5 spikes/s. For cross-correlations, the product of the spontaneous firing rate of both units had to exceed 0.25 spikes/s. The parameter mainly affected by ACh was the cellular average firing rate (Fig. 7). Autocorrelations were affected to a lesser degree and the percentage of affected cross-correlations was very small. These relationships were consistent for all the different groups, except the group of far units recorded with the DC setup, where autocorrelations were rarely affected.

Discussion

The effects of exogenously applied cholinergic agents on cellular activity and functional neuronal connectivity in the auditory cortex of anaesthetized guinea-pigs were studied. In agreement with previous reports, iontophoresis of ACh and of the muscarinic receptor agonist, carbachol, were shown to modulate the ongoing activity of cortical cells. These changes in the firing rates of the cells usually occurred in the absence of a noticeable change in the firing patterns of the neurons, as judged by their autocorrelograms. The effects of ACh and carbachol were studied at the intercellular level by the use of simultaneous recordings of up to 12 units and of cross-correlation techniques. Despite a significant percentage of cells showing a modification in their firing rates upon application of the cholinergic agents, the functional coupling between units, as judged by their cross-correlograms, was almost unchanged. These findings are consistent with the involvement of ACh in neuromodulation of the auditory cortex by modifying the discharge rate of cortical units. They do not support significant promotion by endogenous ACh of synchronization of the firing activity of spontaneously active neurons. However, they do not disprove or support the hypothesis that ACh promotes synchronization of an assembly of neurons upon co-activation by a coherent sensory input (Munk et al., 1996).

Distribution of AChE in the auditory cortex of the guinea-pig

Although various enzymatic histochemical, immunocytochemical and autoradiographic procedures have been used to elucidate the cholinergic innervation of neocortical areas of many species commonly used for experimentation (Bear et al., 1985; Lysakowski et al., 1989), a description of cholinergic innervation in the guinea-pig was not available. In these preparations, the histochemical staining for AChE corresponds well with that for other cholinergic markers (Mesulam and Geula, 1992; Criswell and Brandon, 1993). The distribution of AChE in the auditory cortex of the guinea-pig (Fig. 1) is similar to that previously published for the neocortex of the cat (Bear et al.,
Effects of cholinergic agonists on the spontaneous activity of auditory cells

Methodological considerations

In this study, both the endogenous agonist ACh and the muscarinic agonist carbachol were applied. Of the two agonists, carbachol, because of its lower rate of hydrolysis, is better suited to affect the ongoing activity of neurons situated far from the site of ejection. However, data generated with both agonists was pooled for subsequent analysis, since differences in the qualitative effects induced by the two agonists and in the percentage of cells modified (Table 1) were not significant. This agrees with previous reports that in the
Effect of acetylcholine iontophoresis on the functional coupling of two simultaneously recorded units. The asg values for 0–20 ms time lags between the two units during spontaneous activity periods before and after the application of acetylcholine (ACh; 200 nA, 50 s, vertical dotted lines) are plotted as a function of time (upper panel, each division 100 ms). These spontaneous periods were intermingled with periods in which auditory stimuli were applied, but these periods were excluded from the analysis. The firing rates of both units for the same time periods are also shown (middle and lower panel). Cross- and autocorrelograms (±200 ms, 4 ms bin, computing time 300 s) are shown for selected time periods (horizontal bars below rate meters) before (left), immediately after (middle) and during recovery from (right) an acetylcholine application. The asg values are given in the upper left corner of each histogram. The average (central dashed line) and the confidence lines for $\alpha = 0.01$ (upper and lower dashed lines) were estimated within -200 to -120 ms time lags. For each unit, the quality of unit separation is in the upper right corner of the left histogram.

**Table 4. Dependence on spontaneous initial correlation levels (asg<sub>spnt</sub>) of the sensitivity of functional couplings to cholinergic application**

<table>
<thead>
<tr>
<th>Time lag</th>
<th>asg&lt;sub&gt;spnt&lt;/sub&gt; Neuronal pairs affected</th>
</tr>
</thead>
<tbody>
<tr>
<td>0–20 ms</td>
<td>all 2.8% (47/1670)* 3.5% (53/1497)</td>
</tr>
<tr>
<td>0–300 ms</td>
<td>&gt;0.5 SD 4.0% (23/564) 6.0% (42/695)</td>
</tr>
<tr>
<td></td>
<td>&gt;1 SD 5.5% (17/308) 6.8% (28/408)</td>
</tr>
<tr>
<td></td>
<td>&gt;2 SD 6.7% (8/119) 8.1% (13/161)</td>
</tr>
<tr>
<td></td>
<td>&gt;3 SD 3.0% (2/65) 7.7% (5/65)</td>
</tr>
<tr>
<td></td>
<td>&lt;0.5 SD 2.2% (24/1106) 1.1% (11/1015)</td>
</tr>
</tbody>
</table>

*Percentage of total (affected/total for category).

Excitatory effects of cholinergic agonists

In previous studies, the activation of cholinergic receptors was extensively analysed by intracellular recordings in *in vivo* and *in vitro* preparations. A depolarization of pyramidal cells accompanied by an increase in membrane resistance was reported upon iontophoresis of ACh during intracellular recordings in the anaesthetized cat (Krnjevic et al., 1971). This increase in membrane resistance is due to a blockade of potassium conductances, the most prominent of which are the voltage-dependent M current ($I_M$), the Ca$^{2+}$-dependent $I_{AHP}$

Somatosensory cortex of the unanaesthetized (Basant et al., 1990) and anaesthetized (Lamour et al., 1982) rat, the proportion of cholinceptive cells and the laminar distribution of ACh- and carbachol-induced effects is similar and that 86% of the cells excited by carbachol are also excited by ACh.
current, and the voltage-independent leak current (reviewed in McCormick, 1992a). Blockade of $I_{M}$, and especially of $I_{NH}$, by muscarinic agonists results in reduction of adaptation of spike frequency and an increase in responsiveness to depolarizing inputs. Upon the application of ACh in the cortex, increased excitability is expected because of the biophysical properties of the muscarine sensitive currents. Indeed, most studies, including the present one, of ACh-sensitive cells in the auditory (McKenna et al., 1988; Metherate et al., 1990), visual (Sillito and Kemp, 1983; Sato et al., 1987; Muller and Singer, 1989) and somatosensory (Lamour et al., 1982, 1988; Metherate et al., 1988; Bassant et al., 1990) cortices demonstrate the facilitatory effects of local iontophoretic applications of ACh. However, in these studies the percentage of cells with an increased level of spontaneous activity varied from 16 to 86%, with an average of 39%, which is slightly more than the 25% observed in this study. Differences in the cortical areas, preparations and species used could explain most of the apparent discrepancies and limit the comparability of these studies. For example, in the above studies, 49.3% of the cells showed increased levels of spontaneous activity in awake animals, compared to only 30.3% in anaesthetized animals. The effect of neuroactive substances can be modified by anaesthesia (Krnjevic and Phillis, 1963; Bassant et al., 1990). The difference between the observed effectiveness of ACh in awake and anaesthetized animals could result from the depressive effect of anaesthetics on the average firing rate, since it has been reported that the probability of a neuron being excited by ACh varies according to the level of spontaneous activity (Krnjevic et al., 1971; Jones and Olpe, 1984). More recently, Bassant and co-workers (1990) demonstrated that the probability of cells being excited by ACh increased from 40% for initially silent cells to 70% for cells with spontaneous activity levels of 10—20 spikes/s, but was lower for cells with higher spontaneous activity rates. However, in our study such a relationship between cholinceptive and spontaneously active cells was not observed, which agrees with results obtained in the cat somatosensory cortex (Metherate et al., 1988) and
questions the linkage through the effect on firing rate levels of anesthesia and sensitivity to ACh.

Inhibitory effects of cholinergic agonists

In this and other studies (Randic et al., 1964; McKenna et al., 1988), ACh decreased the spontaneous activity of a relatively high proportion of cells. In neocortical slices of the guinea-pig, an ACh-induced bicuculline-sensitive hyperpolarization, with a reversal potential near the beginning of the application by several tens or hundreds of minutes. This contrasts with the effects of iontophoresed glutamate (Krnjevic and Prince, 1989). Other presynaptic muscarinic actions that reduce excitation (Segal, 1982) could also contribute to the observed inhibition of spontaneous activity.

As in other studies (Sato et al., 1987; Bassant et al., 1990), the effect of ACh and carbachol observed here was usually delayed from the beginning of the application by several tens or hundreds of seconds and could outlast the application period by several tens of minutes. This contrasts with the effects of iontophoresed glutamate (e.g. Haidarliu et al., 1995) and GABA (e.g. Shulz et al., 1993), which usually show sharp onsets and offsets. The delay in the onset of the excitatory action of ACh could be due, at least partially, to the initial cholinergic inhibition (McCormick and Prince, 1986). Also for far cells in our experimental conditions, the latency introduced by the diffusion of the drug could contribute to the delayed response.

In a few cases, biphasic inhibitory and excitatory responses to ACh were recorded in the auditory cortex of the anaesthetized guinea-pig (Fig. 2), similar to observations in the anaesthetized cat (Krnjevic et al., 1971). These biphasic effects are dependent on the site of application of ACh with respect to the soma and dendrites of the cell (Fig. 4 in McCormick and Prince, 1986; Segal, 1982; Markram and Segal, 1992). The hyperpolarizing responses are readily elicited when the applications of ACh are distant from the site of recording, while slow depolarizations are usually seen upon somatic applications of ACh. The biphasic effects observed in this study (Table 2 and Fig. 2) could be the result of a drift of a point of activation due to diffusion of the cholinergic agonist. The dependency of the polarity of the effect on the application site could also explain the large variation between the reported results of different studies with regard to the percentages of excited cells compared with those of inhibited cells. This variation could at least partially be the result of variations in the electrode-to-soma distances, which are caused by different electrode types. For example, glass pipettes usually permit only somatic recordings, while tungsten electrodes permit also distant recordings.

Effects of cholinergic agonists on the burstiness of cortical cells

If neurons behaved as Poisson processes, analysis of average firing rates would be adequate for characterizing the single-cell effects. However, neocortical neurons usually exhibit a serial dependency between their spiking intervals (Abeles, 1982b). One example is cortical neurons that discharge in bursts of several action potentials (Legendy and Salcman, 1985). A typical intrinsic bursting cell (McCormick et al., 1985) can have brief periods of high-frequency activity (120—250 spikes/s). In neocortical cells, as well as hippo-
Campal intrinsic bursting cells, a typical burst is composed of three to five Na+ spikes that ride on a slow-activating Ca2+-dependent depolarization.

A bursting tendency is reflected by a peak in the autocorrelogram (Eggermont et al., 1993). The 'classical' burst, composed of three to five spikes in close temporal contiguity, usually lasts <20—30 ms (McCormick et al., 1985). However, autocorrelograms also exhibit wider peaks of a few hundreds of milliseconds. Such peaks could reflect burst-like phenomena with longer time scales (Legendy and Salcman, 1985) and/or network dynamics (Ginzburg and Sompolinsky, 1994). In order to cover both cases, and to be compatible with the cross-correlation analysis, modifications in autocorrelograms were quantitatively tested by computing the asg values for time lags of both 20 and 300 ms.

In a relatively small number of cases (14 and 16% for the 20 and the 300 ms time lags respectively) the asg values of the autocorrelograms were modified. In the majority of these cases, a decrease in the tendency of the cell to fire in a repetitive way for both time windows was observed (Fig. 3), which agrees with the cholinergic modulation of the neuronal firing mode in the cerebral cortex and thalamus of the guinea-pig in vitro (reviewed in McCormick, 1992a). In the dorsal lateral geniculate nucleus of the guinea-pig (McCormick, 1992b) and in the auditory cortex of the rat (Metherate et al., 1992), application of ACh to rhythmically oscillating neurons to a synchronized to a desynchronized EEG state, a transition that has been traditionally associated with the activation of central cholinergic structures (reviewed in Semb, 1991). Whether pairs of cortical neurons were desynchronized by cholinergic applications was tested in this study using cross-correlations.

**Effects of cholinergic agonists on functional connectivity**

A change in the ongoing activity of a given neuron by ACh could reflect changes in its own excitability level and/or changes in network dynamics or functional connectivity. The cross-correlation technique (Perkel et al., 1967b) is a widely used method for estimating the functional connectivity of simultaneously recorded pairs of neurons (Ahissar et al., 1992a, b; Ahissar and Ahissar, 1994). The adequacy and sensitivity of cross-correlation analysis have been discussed extensively (Aertsen and Gerstein, 1985; Melssen and Epping, 1987) and measures for quantifying interneuronal connections ('efficacy' and 'contribution' in Levick et al., 1972; 'correlation coefficient' in Abeles, 1982b; 'visibility index' in Aertsen and Gerstein, 1985) have been defined. Due to the highly parallel structure of the cortex and the presence of multiple hidden sources of excitatory and inhibitory
connections, it is generally accepted that these quantifications provide an estimation of the effective connectivity (Aertsen et al., 1989) or functional coupling (Aertsen et al., 1994), and not of the structural connectivity, between two simultaneously recorded units. Most of the cross-correlograms computed in this study were of the ‘common input’ type, in agreement with other studies of sensory cortices (Krüger and Aiple, 1988; Vaadia et al., 1991; Nelson et al., 1992).

This type of correlogram can reflect correlated activity over large groups of neurons, a case in which the detection of changes occurring in single synapses is impossible. This study was concerned with detecting changes in coordinated activity within recorded neuronal groups. If ACh affected a significant amount of synapses, the effect should have been reflected in the coordinated activity.

Since few cross-correlograms exhibited a statistically significant change upon application of ACh, the sensitivity of this method for the detection of significant changes was of concern. However, the observed significant changes in the asg or modified cc’ values, the majority of which were in the direction of an increase of the coupling, were usually very strong (>4 SD above background; Figs 4 and 5), indicating that this method was sensitive enough to detect such changes when they occurred. Whether some cholinergic-induced changes in the cross-correlograms were smaller than or equivalent to changes when they occurred. Whether some cholinergic-induced changes in the cross-correlograms were smaller than or equivalent to changes when they occurred. Whether some cholinergic-induced changes in the cross-correlograms were smaller than or equivalent to changes when they occurred.

Furthermore, these excitability changes are not large enough to be reflected in the coordinated activity. This study was supported by the Israel Science Foundation, CNRS, HFSP and the French Ministry of Foreign Affairs during his visits to the Weizmann Institute.

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Abbreviations

\[\begin{align*}
ACh & \quad \text{acetylcholine} \\
AChE & \quad \text{acetylcholinesterase} \\
asg & \quad \text{asynchronous gain} \\
cc’ & \quad \text{modified correlation coefficient} \\
CE & \quad \text{combined electrode} \\
ChAT & \quad \text{choline acetyltransferase} \\
DC & \quad \text{double combined electrode configuration} \\
GABA & \quad \text{γ-aminobutyric acid} \\
SC & \quad \text{single combined electrode configuration} \\
TE & \quad \text{tungsten-in-glass electrode}
\end{align*}\]

References

Cholinergic modulation of correlated activity


