To the Editors:

BISCIONE et al. [1] reported a significant association of upper airway Chlamydia pneumoniae RT-PCR positivity in atopic asthmatics (cumulative rate 22%) compared with nonatopic nonasthmatic spouses (9%). In the discussion, BISCIONE et al. [1] argued in favour of acute, rather than chronic, infection as the explanation for their observation that detection was mostly intermittent rather than persistent. I would like to address three issues raised by this interesting study.

First, in the introduction, BISCIONE et al. [1] stated that C. pneumoniae serology could not differentiate acute from other infections. It is correct to state that serology cannot diagnose chronic infection, but there are established serological criteria for both acute primary and acute secondary infection using the microimmunofluorescence (MIF) test [2]. They also implied that C. pneumoniae serology was nonspecific, i.e. cross-reactive with other Chlamydia species [1], but specificity can be achieved by parallel measurement of other Chlamydia species. For example, the first study they cited as exhibiting these deficiencies was that of HAHN et al. [3], who performed a prospective microbiological and serological study that included acute and convalescent C. pneumoniae MIF serology and was, therefore, able to distinguish serological acute C. pneumoniae infection from other serological patterns. In addition, HAHN et al. [3] included species-specific testing for T. trachomatis antibody as a control, and reported that subjects without evidence for an acute C. pneumoniae infection had a strong, statistically significant and specific “dose-response” association of C. pneumoniae antibody with wheezing and acute asthmatic bronchitis. We interpreted these serological associations as consistent with either reinfection or chronic infection. Serial MIF testing using acknowledged criteria [2] would have established whether the positive detections reported by BISCIONE et al. [1] were related to acute infection or not.

Secondly, BISCIONE et al. [1] stated (correctly in my opinion) that their data could not distinguish acute infection, reactivation, colonisation or chronic infection. It is unlikely that a 22% cumulative incidence rate over 3 months was caused by acute exogenous infections because the annual nonepidemic C. pneumoniae acute infection rate in the adult population is <2% [4]. Acute C. pneumoniae infections can be asymptomatic or associated with only minor respiratory complaints, but a significant minority will cause lower respiratory tract illness [5]. It would be informative to know whether the BISCIONE et al. [1] study was conducted during an epidemic of C. pneumoniae infection in the community, and whether any of the positive detections were associated with an acute respiratory illness.

Thirdly, interpretation of the results is confounded by the mismatch in atopic status introduced by comparing atopic cases with nonatopic controls, i.e. one could argue that atopes are more susceptible to infection than nonatopes, independent of disease status. Future research should include a combination of sensitive nucleic acid detection, serial serological testing, clinical data and appropriate control groups to address the issue of exactly what type of Chlamydia pneumoniae infection is associated with asthma. Uncertainty about the exact type of infection, however, should not delay performance of clinical trials to establish whether asthma is treatable with anti-chlamydial antimicrobials.

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REFERENCES


From the authors:

We thank D.L. Hahn for his interest in our study [1] and for commenting upon the interesting issues he raised relating to the study and its interpretation, particularly regarding the presence of acute infection, reactivation, colonisation or chronic infection. These are important issues and informed debate about them is to be welcomed.

D.L. Hahn comments that, in the introduction, we stated that Chlamydia pneumoniae serology could not differentiate acute from other infections. In fact, we stated that serology cannot reliably differentiate between past and present infection, or acute and chronic infection. We acknowledge that there are published proposed serological criteria for acute primary infection using the microimmunofluorescence (MIF) test [2]. However, these authors stated that “standardized definitions
for ‘acute infection’ and ‘past exposure’ are proposed”, and that “these standards should be applied in future investigations and periodically modified as indicated”. Although certainly the best advice available to date, we believe that these excellent recommendations are certainly not the last word in *C. pneumoniae* diagnosis, which remains a very much still evolving field.

As D.L. Hahn states, we also implied that *C. pneumoniae* serology may be (not “was”) nonspecific, i.e. cross-reactive with other Chlamydia species [1]. We stand by this comment, as there is data indicating this may be the case [3]. We do not wish to imply that all or even most *C. pneumoniae* serology is nonspecific, and we believe that most well-performed MIF testing is indeed specific. However, we do believe that more objective, more specific tests are needed to complement MIF in the serodiagnosis of *C. pneumoniae* infection. Thankfully, the number of such available tests is increasing and they are being used in ongoing studies of *C. pneumoniae* in asthma [4]. We hope that further studies such as these will improve our diagnostic accuracy in due course. We agree with D.L. Hahn that specificity can be improved by parallel measurement of other Chlamydia species and other unrelated organisms, as HAHN et al. [5] and WARK et al. [6] have previously carried out. Such methodological rigour is to be encouraged.

D.L. Hahn states that serial MIF testing using acknowledged criteria [2] would have established whether the positive (PCR) detections we reported [1] were related to acute infection or not. Regrettably, serial blood sampling was not part of our study design, as the original design was conceived to investigate rhinovirus infections [7], for which serology is impractical. The decision to use these samples to investigate the presence of *C. pneumoniae* was only taken once the rhinovirus study was completed. We agree that such serial serological testing would have been desirable and would have added to the evidence, but believe that “established” is too strong a word. The debate regarding *C. pneumoniae* diagnostic methods will certainly continue until clearly accepted gold standard tests are available. Until such time, the data available can only be interpreted in an unbiased manner, and we strongly support D.L. Hahn’s call for future research to include a combination of sensitive nucleic acid detection, serial serological testing and appropriate control groups to address the issue of exactly what type of *C. pneumoniae* infection is associated with asthma and whether it plays a role in disease pathogenesis.

We thank D.L. Hahn for commenting upon our discussion of the detection limit of the PCR assay, or to short-term reactivation.

We also agree with D.L. Hahn that our results do not permit us to conclude whether the increased detections relate to atopic status or asthmatic status, as the study design compared asthmatic cases with nonasthmatic controls. It would clearly be desirable to perform a study including three populations (atopic asthmatic, atopic nonasthmatic and normal nonastomatic subjects), in order to answer this question; however, our spouse-pair design, adopted to control for exposure to infectious agents as closely as possible [7], did not allow this.

Finally, we support D.L. Hahn’s conclusion that uncertainty about the exact type of infection should not delay performance of clinical trials to establish whether asthma is treatable with antichlamydial antimicrobials. These studies should be high-quality, randomised, placebo-controlled studies, and should be supported by further studies on the relationships between *Chlamydia pneumoniae* detection and asthma pathogenesis, to help define which populations are most likely to benefit. We believe such studies should initially be focused on populations that are least well served by currently available asthma treatments, namely, acute asthma exacerbations, new onset adult asthma and severe asthma. It is encouraging to note that, in the context of acute exacerbations, where evidence for a role for *Chlamydia pneumoniae* infection is increasing [6, 4] and where concerns about the induction of antimicrobial resistance to antibiotic therapy would be less worrisome as therapy would be short term, just such a high-quality, randomised, placebo-controlled study is already underway [4].

**REFERENCES**


2 Dowell SF, Peeling RW, Boman J, et al. Standardizing *Chlamydia pneumoniae* assays: recommendations from the
Nasal potentials at high altitude

To the Editors:

In their recent publication in the European Respiratory Journal, SARTORI et al. [1] explain the difference between their results and previous studies by arguing that in other studies “no particular care was taken to locate the electrode in the inferior turbinate”, without contacting the investigators who performed the measurements that they criticise. This argument is not acceptable for us. We can assure them that we paid very careful attention to the placement of the nasal electrode as we are well aware that potentials vary considerably in magnitude in different regions of the nose [2, 3]. The fact that the magnitude of measured potentials is comparable among all studies at high altitude rules out a significant effect of the site of recording and disproves the argument by SARTORI et al. [1].

We would like to make further observations on the discrepant results which are summarised in table 1. In a study by MAIRBAURL et al. [5], performed in 1999 in freezing temperatures and strong winds, subjects reported dryness of the nasal epithelium. This was not a problem in a second study performed in 2003 [6] when weather conditions were warm and nasal dryness was prevented with aerosolised isotonic saline. This manoeuvre entirely prevented the hyperpolarisation of total nasal potential difference. MASON et al. [4] and SARTORI et al. [1] did not observe a problem with nasal dryness, although the subjects in MASON et al. [4] bathed their nostrils with isotonic saline twice daily. These variations indicate the problems and difficulty of interpretation of nasal potential difference measurements. MAIRBAURL et al. [5] found no statistically significant change in the amiloride-sensitive change in the nasal potential difference (NPamil) in high-altitude pulmonary oedema (HAPE)-susceptible subjects; only a nonsignificant trend was reported, whereas, in another study by MAIRBAURL et al. [6], significantly decreased NPamil in HAPE was seen, again pointing to possible effects of nasal dryness. This argument is strengthened by the increase seen in the chloride-sensitive change in the nasal potential difference (NPaci) reported by both MAIRBAURL et al. [5] and MASON et al. [4]. Increased NPaci is compatible with increased nasal secretion. This possibility was not addressed in the study by SARTORI et al. [1].

In summary, these results indicate that the potential across the nasal epithelium might very well be affected by climatic conditions [7] to which the nose is exposed but to which the alveolar epithelium is not [1, 4]. Due to this, particular caution must be exercised when extrapolating data obtained at the nasal epithelium to make claims about changes occurring at the level of the alveolar epithelium.

| TABLE 1 | Studies reporting the change in nasal potential difference upon ascent to high altitude |
|---|---|---|---|---|---|---|
| Change upon ascent to high altitude | Co-NPtot | HAPE-NPtot | Co-NPamil | HAPE-NPamil | Co-NPamil-is | HAPE-NPamil-is |
| SARTORI et al. [1] | NS | # | NS | NS | # | # |
| MASON et al. [4] | * | * | NS | NS | * | * |
| MAIRBAURL et al. [5] | * | * | # | NS | * | * |
| MAIRBAURL et al. [6] | # | # | # | # | # | # |

Co: controls; NPtot: total nasal potential difference; HAPE: high-altitude pulmonary oedema; NPamil: amiloride-sensitive change in the nasal potential difference; NPamil-is: amiloride-insensitive nasal potential difference; #: decreased potential difference, more positive values; *: increased potential difference, more negative values; NS: no significant change. Symbols are shown only when changes were reported to be statistically significant.