Different pH requirements are associated with divergent inhibitory effects of chloroquine on human and avian influenza A viruses
Livia Di Trani†1, Andrea Savarino*†2, Laura Campitelli2, Sandro Norelli2, Simona Puzelli2, Daniela D'Ostilio2, Edoardo Vignolo1, Isabella Donatelli2 and Antonio Cassone2

Address: 1Dept. of Food and Animal Health, Istituto Superiore di Sanità, Rome, Italy and 2Dept. of Infectious, Parasitic, and Immune-mediated Diseases, Istituto Superiore di Sanità, Rome, Italy

Email: Livia Di Trani - livia.ditrani@iss.it; Andrea Savarino* - asavarino@medscape.com; Laura Campitelli - laura.campitelli@iss.it; Sandro Norelli - xsandro76@hotmail.com; Simona Puzelli - simona.puzelli@iss.it; Daniela D'Ostilio - danieladostilio@hotmail.it; Edoardo Vignolo - edoardo.vignolo@iss.it; Isabella Donatelli - isabella.donatelli@iss.it; Antonio Cassone - antonio.cassone@iss.it
* Corresponding author †Equal contributors

Abstract
Chloroquine is a 4-aminoquinoline previously used in malaria therapy and now becoming an emerging investigational antiviral drug due to its broad spectrum of antiviral activities. To explore whether the low pH-dependency of influenza A viruses might affect the antiviral effects of chloroquine at clinically achievable concentrations, we tested the antiviral effects of this drug on selected human and avian viruses belonging to different subtypes and displaying different pH requirements. Results showed a correlation between the responses to chloroquine and NH4Cl, a lysosomotropic agent known to increase the pH of intracellular vesicles. Time-of-addition experiments showed that the inhibitory effect of chloroquine was maximal when the drug had been added at the time of infection and was lost after 2 h post-infection. This timing approximately corresponds to that of virus/cell fusion. Moreover, there was a clear correlation between the EC50 of chloroquine in vitro and the electrostatic potential of the HA subunit (HA2) mediating the virus/cell fusion process. Overall, the present study highlights the critical importance of a host cell factor such as intravesicular pH in determining the anti-influenza activity of chloroquine and other lysosomotropic agents.

Background
A second look at selected compounds is giving new life to several abandoned therapies and new applications for existing drugs [1-3]. One such example is provided by chloroquine, being dismissed from antimalarial treatment and finding new applications in the clinical management of autoimmune diseases, tumours and non-malarial infections [4,5]. The use of chloroquine in the clinical management of a viral infection was first considered in the 1990s, on the basis of its effects on HIV-1 [6,7]. The drug is now being tested as an investigational antiretroviral [8].

Some of us previously analysed the reported effects of chloroquine on replication of several viruses and concluded that the drug should be studied as a broad spectrum antiviral agent against emerging viral infections, being relatively well tolerated, cheap, and immediately available.
available worldwide [9]. As a weak base capable of accumulating within cellular organelles, chloroquine appears to be capable of interfering with pH-dependent steps in the replication of several viruses. Other mechanisms of viral inhibition by chloroquine, such as inhibition of polynucleotidyl transferases have, however, been considered [7]. In 2003–2005, chloroquine was studied as a promising in vitro anti-SARS agent [9-11] and recently entered clinical trials against chikungunya fever [12].

The broad-spectrum antiviral effects of chloroquine deserve particular attention in a time in which there are several cases of avian influenza A virus transmission to humans from poultry, and the availability of antiviral drugs is fundamental during preparation and evaluation of effective vaccines. Chloroquine inhibition of both type A and B influenza viruses was first described in the 1980s [13,14]. The concentrations employed in these studies were however too high to allow a theoretical transposition to in-vivo settings. Anecdotal reports of clinical benefits derived from a related compound, i.e. quinine, date back to the Spanish influenza pandemic of 1918/19. However, it was not until last year that the anti-influenza virus effects of chloroquine at clinically achievable concentrations were studied, in view of a possible application of this drug in the clinical management of influenza [4,15]. Investigations still have to be done on this topic. For example, the mechanisms of orthomyxovirus inhibition by chloroquine have been uncertain at the clinically achievable concentrations adopted in the most recent studies [4,15], as well as the effects of chloroquine on field isolates, including avian strains potentially transmittable to humans.

We here report the results of an initial evaluation of susceptibility to chloroquine of human and avian influenza A viruses. Susceptibility to chloroquine appears to be dependent on the pH requirements of the viruses and the electrostatic potential of haemagglutinin subunit 2 (HA2), which is involved in virus/cell fusion. Accordingly, the antiviral effects are exerted at an early step of virus replication.

**Results**

We first tested the effects of chloroquine on low-pathogenic (LP) A/Ck/Il/9097/97 (H5N9) virus, isolated from poultry in Italy. We found that chloroquine dose-dependently inhibited the viral cytopathic effect with a 50% effective concentration (EC50) of 14.38 μM, in cells infected with the H5N9 virus at approx. 10^4 50% tissue culture infecting doses (TCID50)/ml (Fig. 1a). Although this value was rather high, some of the inhibitory concentrations matched the blood concentrations reported in individuals under acute antimalarial treatment (1–15 μM). The inhibitory effects were confirmed using quantitative reverse transcriptase real-time PCR (qRRT-PCR) (Fig. 1b).

Ooi et al. (2006) [15] recently reported that chloroquine inhibited human H3N2 and H1N1 viruses with EC50 values in the range of 0.84 – 3.60 μM. To investigate whether the discrepancies with the inhibitory values reported above were due to the type of virus or to the different conditions and methods adopted, we tested the effects of chloroquine on replication of recent human H3N2 and H1N1 viral isolates under conditions similar to those adopted for the H5N9 avian influenza virus. Using the test of inhibition of viral cytopathogenicity, we found that chloroquine inhibited the H3N2 virus (10^4 TCID50/ml) with an EC50 of 1.53 μM (Fig.1c). Inhibition was confirmed using qRRT-PCR, both under similar conditions and at lower MOIs (Fig.1d), thus confirming the results of Ooi et al. (the assay adopted by these authors employs a lower MOI than routinely used by our group). Results obtained with H1N1 viruses (10^4 TCID50/ml) showed a similar drug susceptibility for the human strain (EC50 = 1.26 μM), in full agreement with Ooi et al. [15], but no response to clinically achievable drug concentrations in an avian strain (IC50 > 20 μM; data not shown). These data suggest a more pronounced inhibitory effect of chloroquine on human H3N2 and H1N1 viruses than on avian H5N9 virus replication.

Since 1) chloroquine is thought to interfere with pH-dependent steps of the life cycles of several viruses [9], and 2) some of us reported different pH requirements in influenza A viruses infecting different avian species [15], we investigated whether response of influenza viruses to chloroquine might depend on the different pH requirements of the human and avian viruses.

Thus, we analysed the response of human H3N2 virus (good chloroquine responder) and avian H5N9 virus (poor chloroquine responder) to ammonium chloride (NH4Cl; 40 mM), a lysosomotropic agent known to increase the pH of intracellular vesicles. Results showed a good response of the chloroquine-sensitive H3N2 virus to NH4Cl inhibition of viral cytopathogenicity (100% inhibition under conditions described above). Conversely, the lower chloroquine-sensitivity of H5N9 virus was associated with lack of response to NH4Cl (data not shown). This observation raised the hypothesis that cellular pH might be a critical factor for chloroquine inhibition of influenza virus.

To explore this possibility, the action of chloroquine was tested on two avian H7N3 viruses whose haemagglutinins (HAs) differ in two amino acid positions (i.e. residue 261 in the HA1 subunit and residue 161 in HA2, the latter being the HA subunit mediating the fusion process), and
which display different pH requirements [16]. The two viral strains showed a marked discrepancy in the response to chloroquine. In particular, A/Mallard/It/43/01 (H7N3) virus, which had been shown to be relatively more independent from pH increase than A/Ty/It/220158/02 virus [16], was insensitive to clinically achievable chloroquine concentrations (EC<sub>50</sub> > 20 μM). In contrast, chloroquine exerted some inhibitory effect on A/Ty/It/220158/02 replication (EC<sub>50</sub> = 14.39 μM), although the response of this virus to chloroquine was lower than that of H3N2 virus.

The different pH requirements of the two viruses were confirmed by the different responses to NH₄Cl (0% inhibition for A/Mallard/It/43/01; 30% inhibition for A/Ty/It/220158/02). This result raises the hypothesis that chloroquine inhibits pH-dependent events involving HA.

To further explore this possibility, the isoelectric point was calculated for HAs of all viruses used in the present study, and the electrostatic potential was mapped on the protein surfaces of 3D models obtained by homology...
with the crystal structures of matched subtype representatives. Results showed a negative correlation between the isoelectric point of HA2 and the IC_{50} of chloroquine in vitro (r = -0.87; P = 0.024; Fig. 2). Instead, no correlation was found with the isoelectric point of HA1 (P > 0.05; data not shown). Although the viruses studied here belonged to different subtypes, chloroquine-resistant viruses, independently of the subtype, showed a more marked negative surface potential of HA2 than chloroquine-sensitive viruses (Fig. 2). Viruses with intermediate drug sensitivity showed intermediate characteristics (Fig. 2). We conclude that structural determinants in HA2 are associated with response of influenza A viruses to chloroquine.

If the hypothesis of a pH dependent inhibitory action of chloroquine was correct, the timing of drug inhibition should match that of virus/cell fusion, an early step of virus replication occurring in endosomes and requiring a low pH (approx. pH 5–5.5) in several, but not all, influenza A viruses, as shown by previous studies [17]. As the assays for detection of antiviral effects adopted in the first

Figure 2
Correlation between electric characteristics of haemagglutinin subunit 2 (HA2) and response to chloroquine of influenza A viruses. A) Correlation between EC_{50} of chloroquine (CQ) on viral cytopathogenicity (presented as Log values, x axis) and isoelectric point of HA2 (pH value at which the protein is neutral; y axis). The line best fitting the data points is shown. Isoelectric points were calculated based on the protein sequence using the web interface in Ref. [35]. B-G) Theoretical three-dimensional models for HA2 subunits of the viruses adopted in the present study, shown in ranked order of sensitivity to chloroquine (from resistant to clinically achievable concentrations to fully sensitive). B) A/Mallard/Italy/43/01 (H7N3); C) A/ Mallard/Italy/66/96 (H1N1); D) A/Chicken/Italy/9097/97 (H5N9); E) A/Turkey/Italy/220158/02 (H7N3); F) Human influenza A/ Panama/2007/99-like (H3N2); G) A/New Caledonia/20/99-like (H1N1). The electrostatic potential is mapped to the protein surface as a range of colours from red (-1.8 V) to blue (+1.8 V). Equipotential surfaces are shown as dotted line grids. Protein sequence data regarding the H7N3 and H5N9 viruses can be found in GenBank with the following accession numbers: AAT37405, AAT37404, AAG60349.
part of this study were designed to allow multiple cycles of viral replication, we designed time-of-addition experiments using the chloroquine-sensitive human H3N2 virus. Chloroquine was added during virus adsorption onto cells (i.e. time 0; T0) and/or at 1, 2, 3 and 4 h post-infection (T1–4). Using qRRT-PCR, we found that the inhibitory effect of chloroquine was highest when the drug was added at T0 (inhibition of viral replication corresponding to 89,36%) and at T1 (inhibition of replication corresponding to 15,53%), whereas the inhibitory activity was completely lost at T2.

If this timing was correct, chloroquine should inhibit influenza A replication by a novel mechanism, and therefore exert additive effects in combination with oseltamivir, inhibiting neuraminidase activity at the late stages of viral replication cycle. To test this hypothesis, human H3N2 virus-infected cells were treated with different chloroquine concentrations in the presence or absence of oseltamivir (10 nM). The virus-infected cells were also incubated with oseltamivir alone, EC50 = 20 nM. Isobologram analysis showed that the two drugs exerted an additive effect (sum of FICs = 1) (data not shown). This result provides further evidence that chloroquine inhibits viral replication by a mechanism different from that of one major anti-influenza drug.

Discussion

The results so far obtained suggest that chloroquine inhibits the replication of those influenza A viruses requiring low pH for proper fusion activation and that the antiviral effects occur at an early stage of viral replication. Supporting evidence comes from: 1) the more sustained inhibitory effect of chloroquine on those viruses whose haemagglutinins (HAs) were found to require low pH for their fusion activity, 2) common HA2 characteristics such as surface potential and isoelectric point in chloroquine-sensitive viruses, and 3) time-of-addition experiments with H3N2 virus.

Chloroquine was found to inhibit a number of cellular processes, some of which do not depend on low pH but might anyway interfere with viral replication. For example, the drug was found to inhibit viral nucleotidyl transferases such as HIV-1 integrase [7]. If chloroquine inhibited influenza A RNA-dependent RNA polymerase, the timing of viral inhibition would not be consistent with that observed in the present study, because RNA replication occurs in the nucleus at later stages [18].

Based on bioinformatic studies, it was recently hypothesized that chloroquine might inhibit UDP-N acetylglucosamine transferase [1], a limiting enzyme in sialic acid synthesis. This specific issue has not been addressed here. Nonetheless, if the antiviral effect of chloroquine reported here were due to inhibition of UDP-N acetylglucosamine transferase, the drug should likely have antagonized the antiviral effect of the neuraminidase inhibitor oseltamivir (acting on detachment of sialic acid-bound virions from parent cells), rendering oseltamivir inhibition unnecessary. Instead, chloroquine was found in the present study to exert antiviral effects that were additive to those of oseltamivir.

Chloroquine is a weak base that is known to affect acid vesicles leading to dysfunction of several proteins [9], and has been shown to inhibit different viruses requiring a pH-dependent step for entry [19–22]. The results of the time-of-addition experiments, performed in the present study using a recent epidemic isolate of human H3N2 virus, are consistent with chloroquine inhibition of pH-dependent steps occurring at an early phase of influenza A virus replication.

Most influenza viruses enter target cells by fusion of the viral and cell membranes at the endosomal pH (approx. pH 5–5.5), although some virus variants can replicate well also at higher pH values [16,17,23]. In a previous paper, the differential growth sensitivity of two naturally occurring variants of an H7N3 virus, isolated from mallards and turkeys, to increased pH values was shown to correlate with different fusion properties [16]. Since chloroquine appears to mimic the effects observed on these two viruses when pH is increased, our data support the hypothesis that the step of virus replication inhibited by clinically relevant chloroquine concentrations is the low-pH dependent haemagglutinin-mediated virus/cell fusion, in agreement with the evidence obtained by other authors at much higher concentrations than those used in this study [13,14]. The correspondence between antiviral effects and isoelectric point of HA2 (i.e. the HA subunit mediating the fusion process) is also consistent with this mechanism. Acidic pH in the endosomal compartment also activates the influenza virus ion channel, M2, that promotes the uncoating of influenza virus in endosomes. However, M2 involvement as a possible target of the antiviral effect of chloroquine is unlikely, since no aminoacid differences were observed in M2 trans-membrane region (the ion-channel domain) between the two avian H7N3 viruses with different chloroquine sensitivities, as already reported [16].

Although a comprehensive study on the variation of fusion pH requirements of influenza A viruses of all HA subtypes and isolates from different hosts is not available, several authors have documented that the threshold pH, at which the HA conformational change and virus-cell fusion occur, is strain-specific [24–26]. Interestingly, the viruses showing highest chloroquine sensitivity also displayed the highest HA2 isoelectric points. Thus, a relation-
ship between isoelectric point and response to pH is apparent. However, a broad study relating the surface electrostatic potential with inactivation by pH would be required to analyse the molecular details of the HA/pH interplay. Analyses of virus production before and after exposure to chloroquine and of the possible changes in HA2 surface potential in viruses rendered resistant to chloroquine after long exposure to the drug will also be necessary.

**Conclusion**

Although association between variables cannot be considered to be equivalent to causation, the results of the present study strongly suggest that pH critically determines the antiviral activity of chloroquine by regulating virus/host cell interactions. The potential use of this compound as an antiinfluenza drug should take into consideration the possibility that even within the same subtype, different strains may present significantly divergent sensitivities to chloroquine as a consequence of their different pH requirements. Moreover, sensitivity to chloroquine may vary in different cell populations susceptible to influenza A virus infection, depending on different capabilities of endosome acidification. Mutations affecting the electrostatic potential of the the HA2 protein subunit of various isolates of the same virus could also be relevant. All these factors should be carefully evaluated when hypothesising a potential clinical utilisation of chloroquine against influenza A viruses.

**Materials and methods**

**Cells and virus stocks**

Madin Darby Canine Kidney (MDCK) cells were obtained from the American Type Culture Collection. The following viruses were used in this study: two recent human strains, A/Panama/2007/99-like (H3N2) and A/New Caledonia/20/99-like (H1N1), and four LP avian influenza viruses, A/Chicken/Italy/9097/97 (H5N9), A/Turkey/Italy/220158/02 (H7N3) and A/Mallard/Italy/43/01 (H7N3), A/Mallard/Italy/66/96 (H1N1). Virus titration was performed by 50% tissue culture infectious dose (TCID50) in MDCK cells, as described [27], and virus stocks were aliquoted and stored at -70°C until used. All the viruses were from the Istituto Superiore di Sanità (ISS) repository. Virus infection of MDCK cell monolayers was carried out according to standard procedures [28].

**Compounds**

Chloroquine phosphate (7-chloro-4- [4-(diethylamino)-1-methylbutyl]amino]quinoline phosphate, (Sigma) and oseltamivir, a kind gift from Roche was used as a positive control.

**Virus yield assay**

After 24 hours of incubation of the virus-infected cells with different concentrations of the test compounds, under the appropriate conditions, pooled aliquots of the supernatants containing free viruses were subjected to RNA extraction and qRRT-PCR.

**Quantitative Real Time RT-PCR (qRRT-PCR)**

A one-step qRRT-PCR assay was employed, which makes use of minor groove binder (MGB) probe technology, as previously described [29]. Briefly, viral RNA was extracted from infected cell supernatants (QIAmp Viral RNA Mini kit – Qiagen, GmbH, Hilden, Germany) and amplified by RRT-PCR using primers and probe targeting a highly conserved region of the matrix gene of influenza type A viruses. The influenza matrix RNA was also in vitro transcribed from the corresponding DNA template, cloned into a plasmid vector as previously described [29] and used as standard RNA to generate standard curves for quantification of the vRNA in cell supernatants.

**Assay for measurement of antiviral activity**

100 μl of 1.5 × 10⁵ cells/ml MDCK cells in growth medium was seeded into each well in the 96-well microtiter plate. When the cell monolayer was confluent, the culture medium was removed and cells were washed twice with serum-free MEM. Then, 100 μl of the Type A influenza viruses under study containing 10³– 10⁴ TCID₅₀ were inoculated in wells and the plates incubated for 1 hour at 37°C in humidified air of 5% CO₂. The viral suspension was removed and the cells were washed two times; fresh medium containing TPCK-trypsin and chloroquine at different concentrations or NH₄Cl (40 mM) was then added to culture wells in triplicate. Antiviral activity and cytotoxicity measurements were based on the viability of cells that had been infected or mock infected with influenza viruses in the presence of various concentrations of the test compounds. One to three days, depending on the kinetics of cytopathogenicity, after infection the number of viable cells was quantified by a tetrazolium-salt-based-colorimetric method (CellTiter 96 AQueous One Solution kit, Promega, The Netherlands).

**Assessment of the effects of two drugs in combination**

To measure the anti-influenza effects of chloroquine/oseltamivir drug combinations, cell pellets were resuspended in media containing increasing concentrations of the antimalarial in the presence or absence of oseltamivir. A fractional inhibitory concentration (FIC) was then calculated as the ratio: 50% effective concentration (EC₅₀) of drug A in combination with drug B/EC₅₀ of drug A alone. The effect was considered to be additive when the sum of FICs was between 0.8 and 1.2, as previously described [8].
Time-of-addition assay
Monolayers of MDCK cells in 96-well plates were infected with 100 μl of medium containing approximately 10^4 TCID_{50} of H3N2 subtype. After 1 hour of adsorption, cell monolayers were washed twice with serum-free MEM and incubated in fresh medium containing TPCK-trypsin and chloroquine at a concentration of 10 μM. Chloroquine was added at the time of infection or at four different time points thereafter. Eight hours post-infection, a time point at which all progeny virus in the supernatants is derived from the first replication cycle, cell supernatants were collected, viral RNA was extracted and the antiviral activity was determined by using the qRRT-PCR described above.

Viral RNA sequencing
Hemagglutinin genes of H3N2 and H1N1 viruses were sequenced using gene-specific primers, as previously described [30]. Sequence data so far unpublished will be deposited in GenBank by the time of publication of the present article.

Molecular modelling
Three-dimensional models for the HAs of the viruses used in the present study were obtained by homology modelling, using the SWISS model web server [31,32], using, as templates, structures of matched subtype representatives deposited in the protein data bank (PDB) [33]. Hydrogens were added using VEGA-ZZ (University of Milan, Italy) [34], and the structures were then visualised using the Swiss PDB Viewer (SPDBV) program (Swiss Institute of Bioinformatics) [31].

The Coulomb potential was mapped to the protein surface by use of SPDBV using the default relative dielectric constant (solvent = water) of 80. Further information on the algorithm adopted by the program is available in the detailed online description of the program [31].

Authors’ contributions
LDT developed the virus detection assays and participated in the experimental design and manuscript drafting. AS participated in the general design of the study, data analysis and molecular modelling, and drafted the manuscript. LC participated in the experimental design, data analysis and manuscript drafting. SP prepared and sequenced the viral samples and was involved in the assays for antiviral activity. EV, SN and DDO conducted the assays for antiviral activity. ID coordinated the virological activities and participated in the experimental design. AC conceived and coordinated the entire study.

Acknowledgements
The authors are thankful to Roche Italia for providing oseltamivir, and to Dr. Roberto Cauda, M.D., Ph.D., Universitá Cattolica del Sacro Cuore, Rome, Italy, for his advocacy on research on chloroquine as an antiviral.

The research was supported by a FIRB grant from the Italian Ministry of University and Research.

References
1. Cassone A, Cauda R: HIV proteinase inhibitors: do they really work against Candida in a clinical setting? Trends Microbiol 2002, 10:177-178.
2. Jenwitheesuk E, Samudrala R: Identifying inhibitors of the SARS coronavirus proteinase. Bioorg Med Chem Lett 2003, 13:3989-3992.
3. Savarino A: Expanding the frontiers of existing antiviral drugs: possible effects of HIV-1 protease inhibitors against SARS and avian influenza. J Clin Virol 2005, 34:170-178.
4. Savarino A, Di Trani L, Donatelli I, Cauda R, Cassone A: New insights into the antiviral effects of chloroquine. Lancet Infect Dis 2006, 6:67-69.
5. Savarino A, Lucia MB, Giordano F, Cauda R: Risks and benefits of chloroquine use in antirickettsial treatments. Lancet Oncol 2006, 7:792-793.
6. Tsai WP, Nara PL, Kung HF, Oroszlan S: Inhibition of human immunodeficiency virus infectivity by chloroquine. AIDS Res Hum Retroviruses 1990, 6:69-72.
7. Savarino A: A historical sketch of the discovery and development of HIV-1 integrase inhibitors. Expert Opin Investig Drugs 2006, 15:1507-1522.
8. Savarino A, Lucia MB, ter Heine R, Rastrelli E, Rutella S, Majoori G, Huitema A, Boelaert JR, Cauda R: Quinoline antimalarials as investigational drugs for HIV-1/AIDS: in vitro effects on HIV-1 replication, HIV-1 response to antiretroviral drugs, and intracellular antiretroviral drug concentrations. Drug Develop Res 2006, 67:806-17.
9. Savarino A, Boelaert JR, Cassone A, Majoori G, Cauda R: Effects of chloroquine on viral infections: an old drug against today’s diseases? Lancet Infect Dis 2003, 3:722-727.
10. Keyaerts E, Vigen L, Maes P, Neyts J, Van Ranst M: In vitro inhibition of severe acute respiratory syndrome coronavirus by chloroquine. Biochem Biophys Res Commun 2004, 333:264-268.
11. Vincent MJ, Bergeron E, Benjannet S, Erickson BR, Rollin PE, Ksiazek TG, Seidah NG, Nichol ST: Chloroquine is a potent inhibitor of SARS coronavirus infection and spread. Virol J 2005, 2:69.
12. Agence Française de Sécurité Sanitaire des Produits de Santé [http://agmed.sante.gouv.fr/pdf/10/chicungu.pdf]. Press release. April 6, 2006.
13. Yoshimura A, Kuroda K, Kawasaki K, Yamashita T, Maeda T, Ohnishi S: Infectious cell entry mechanism of influenza virus. J Virol 1982, 43:284-293.
14. Shibata M, Aoki H, Tsurumi T, Sugiyama Y, Nishiyama Y, Suzuki S, Maeno K: Mechanism of uncoating of influenza B virus in MDCK cells: action of chloroquine. J Gen Virol 1983, 64:1149-1156.
15. Ooi EE, Chew JS, Loh JP, Chua RC: In vitro inhibition of human influenza A virus replication by chloroquine. Virol J 2006, 3:39.
16. Giannecchini S, Campitelli L, Calzoletti L, De Marco MA, Azzi A, Donatelli I: Comparison of in vitro replication features of H7N3 influenza viruses from wild ducks and turkeys: potential implications for interspecies transmission. J Gen Virol 2006, 87:171-175.
17. Scholtissek C: Stability of infectious influenza A viruses at low pH and at elevated temperature. Vaccine 1985, 3:215-218.
18. Krug RM, Firelli VAC, Jalkunen I, Kazge MG: Expression and replication of the Influenza Virus Genome. In The influenza viruses Edited by: Krug RM. Plenum Press, New York; 1989:89-152.
19. Gonzalez-Dunia D, Cubitt B, de la Torre JC: Mechanism of Borna disease virus entry into cells. J Virol 1998, 72:783-788.
20. Ros C, Burckhardt CJ, Kempf C: Cytoplasmic trafficking of minute virus of mice: low-pH requirement, routing to late endosomes, and proteasome interaction. J Virol 2002, 76:12634-12645.
21. Diaz-Griffero F, Hoschander SA, Brojatsch J: Endocytosis is a critical step in entry of subgroup B avian leukosis viruses. J Virol 2002, 76:12866-76.
22. Bishop NE: Examination of potential inhibitors of hepatitis A virus uncoating. Intervirology 1998, 41:261-271.
23. Daniels RS, Downie JC, Hay AJ, Knossow M, Skehel JJ, Wang ML, Wiley DC: Fusion mutants of the influenza virus hemagglutinin glycoprotein. Cell 1985, 40:431-439.
24. Maeda T, Ohnishi S: Activation of influenza virus by acidic media causes hemolysis and fusion of erythrocytes. FEBS Lett 1980, 122:283-287.
25. Huang RT, Rott R, Klenk HD: Influenza viruses cause hemolysis and fusion of cells. Virology 1981, 110:243-247.
26. Skehel JJ, Bayley PM, Brown EB, Martin SR, Waterfield MD, White JM, Wilson IA, Wiley DC: Changes in the conformation of influenza virus hemagglutinin at the pH optimum of virus-mediated membrane fusion. Proc Natl Acad Sci USA 1982, 79:968-972.
27. Rowe T, Abernathy RA, Hu-Primmer J, Thompson WW, Lu X, Lim W, Fukuda K, Cox Nj, Katz JM: Detection of antibody to avian influenza A (H5N1) virus in human serum by using a combination of serologic assays. J Clin Microbiol 1999, 37:937-943.
28. WHO: Concepts and procedures for laboratory-based influenza surveillance. U.S Department of Health and human services, Centers for disease control, Atlanta, Georgia; 1982.
29. Di Trani L, Bedini B, Donatelli I, Campitelli L, Chiappini B, De Marco MA, Delogu M, Buonavoglia C, Vaccari G: A sensitive one-step real-time PCR for detection of avian influenza viruses using a MGB probe and an internal positive control. BMC Infect Dis 2006, 6:87.
30. Campitelli L, Mogavero E, De Marco MA, Delogu M, Puzelli S, Frezza F, Facchini M, Chiappponi C, Foni E, Cordioli P, Webby R, Barigazzi G, Webster RG, Donatelli I: Interspecies transmission of an H7N3 influenza virus from wild birds to intensively reared domestic poultry in Italy. Virology 2004, 323:24-36.
31. ExPASy Proteomics Server [http://www.expasy.org]
32. Schwede T, Kopp J, Guex N, Peitsch MC: SWISS-MODEL: an automated protein homology-modeling server. Nucleic Acids Research 2003, 31:3381-3385.
33. RCSB Protein Data Bank [http://www.rcsb.org/pdb/home/home.do]
34. VEGA ZZ Homepage [http://www.ddl.unimi.it/vega/index2.htm]
35. Palabra! Isoelectric Point Estimator/Calculator [http://www.nihilnovus.com/Palabra.html]