Animal viruses, bacteria, and cancer: a brief commentary

Jimmy T. Efird1*, Stephen W. Davies2, Wesley T. O’Neal3 and Ethan J. Anderson4

1 Department of Public Health, Brody School of Medicine, East Carolina University, Greenville, NC, USA
2 Department of General Surgery, University of Virginia School of Medicine, Charlottesville, VA, USA
3 Department of Internal Medicine, Wake Forest University School of Medicine, Winston-Salem, NC, USA
4 Department of Pharmacology and Toxicology, Brody School of Medicine, East Carolina University, Greenville, NC, USA

Editted by:
Faith G. Davis, University of Alberta, Canada

Reviewed by:
Li Tang, St. Jude Children’s Research Hospital, USA
Amy Millikan Bell, US Army Public Health Command, USA

*Correspondence:
Jimmy T. Efird, Department of Public Health, Brody School of Medicine, East Carolina University, Greenville, NC 27834, USA
email: jimmy.efird@stanfordalumni.org

Animal viruses and bacteria are ubiquitous in the environment. However, little is known about their mode of transmission and etiologic role in human cancers, especially among high-risk groups (e.g., farmers, veterinarians, poultry plant workers, pet owners, and infants). Many factors may affect the survival, transmissibility, and carcinogenicity of these agents, depending on the animal-host environment, hygiene practices, climate, travel, herd immunity, and cultural differences in food consumption and preparation. Seasonal variations in immune function also may increase host susceptibility at certain times of the year. The lack of objective measures, inconsistent study designs, and sources of epidemiologic bias (e.g., residual confounding, recall bias, and non-randomized patient selection) are some of the factors that complicate a clear understanding of this subject.

Keywords: animal viruses, bacteria, epidemiology, cancer, infection

INTRODUCTION

Humans and animals have coexisted since the beginning of time, sharing viruses, bacteria, and perhaps the etiology of cancers. Approximately 75% of viruses and 50% of bacteria known to cause disease in humans are zoonotic and can be transmitted between animals and people (1). While evolution has provided adaptive immunity against microbes and cancer, the ability to defend against infection is sometimes absent or compromised. Excluding ionizing radiation, sunlight, and tobacco, infection represents the main known cause of human cancers throughout the world. The list is long including cancers of the anogenital track (e.g., residual confounding, recall bias, and non-randomized patient selection) are some of the factors that complicate a clear understanding of this subject.

Keywords: animal viruses, bacteria, epidemiology, cancer, infection

BACTERIA

Animal bacteria also have been implicated in cancer. The occurrence of gliomas in the brain of fowl has been noted in several reports (17–19) and these tumors have been described as having their pathognomonic encephalitic features of a pleomorphic parasite infection (e.g., hypertrophy and hyperplasia of blood-vessels; perivascular infiltration by lymphocytes, plasma cells, and monocytes; and the presence of A-D bodies) (20). Chickens spontaneously and experimentally infected with toxoplasma have been
observed to develop glioma-like tumors (21, 22). A study of 16 human brain tumors observed bodies indistinguishable from the C and D phases of the fowl parasite (23). Epizootic outbreaks of toxoplasmosis have been reported in various avian species and mammals (22, 24, 25). Furthermore, toxoplasma antibodies have been isolated in the blood of exposed sheep farmers, flock animals, herder dogs, mice, and rats (26). Potential cellular mechanisms by which animal viruses and bacteria lead to tumorgenesis are shown in Figure 1.

AGE OF EXPOSURE
Early age at first exposure is critical for the development of many malignancies (27). Fetal or early childhood exposure to infectious agents occurs at a vulnerable time when the immune system is less developed and possibly more susceptible to tumor formation (28). The unrecognized persistence of an oncovirus or microbe in the host increases the chance for this agent to alter the cell cycle and immortalize cells (2, 29). Inoculation of immunologic immature neonate mice with polyoma virus results in tumor formation at multiple sites. In contrast, older mice do not develop tumors in response to polyoma virus either in the laboratory or by natural infection (13). The selective sensitivity of the fetus is further demonstrated in a study of patas monkeys wherein the administration of known carcinogenic agents in the fetal stage cause more tumors than an equivalent dose administered to juvenile monkeys (27, 30). Furthermore, few tumors developed when treatment was delayed until the start of the second trimester, suggesting that sensitivity was greatest during the first trimester.

TRANSMISSION
EXPOSURE ROUTES AND FACTORS INFLUENCING TRANSMISSION
Human exposure may occur in many ways – preparing and consuming animal products, washing with, and drinking well water contaminated with animal fecal coliform, animal bites/scratches, and working in occupations involving regular contact with animals, manure, soil, and/or by-products (e.g., farmers, slaughtering plant workers). Even living down-wind of a farm field fertilized with animal manure poses a potential risk. A list of major sources and exposure routes of animal-to-people transmission of viruses and bacteria is shown in Figure 2. Factors influencing the probability of disease transmission involve the proximity and temporal contact with the infectious organism, length of time that the infectious agent is present, virulence of the agent, incubation period, stability of the agent under varying environmental conditions, population density of carrier animals, husbandry practices, and control of wild rodents and insects (31). The type and maintenance of animal housing also may affect the extent to which individuals working in or around such facilities are exposure to zoonotic viruses and bacteria. Often, animal containment structures (e.g., hen houses, pig pens, cattle barns, and horse stables) may be inadequately ventilated and/or have poor waste removal systems, increasing the exposure of animals and their caretakers to dust, fecal matter, and microbes (32).

MECHANICAL INTERMEDIATE HOSTS
Infectious agents may be transmitted to humans by direct contact, fomite or mechanical vector, or intermediate hosts in which the agent multiples or develops before transmission to animal or human (i.e., metazoonoses). Examples of infectious agents requiring an incubation period prior to transmission include arboviruses, plague, and schistosomiasis (31). In the case of toxoplasmosis infection, contaminated soil and water represent a key source of infection emanating from an intermediate host (33, 34). Indoor/outdoor cats are a significant carrier/transmitter of Toxoplasma, shedding the organism in its feces (34). Oocysts from Toxoplasma gondii also may be transported by cockroaches and...
other bugs and deposited onto food and later consumed by animals and humans (35). In a recent study, eating raw oysters, clams, or mussels was identified as a new risk factor for *T. gondii* infection (36). The *T. gondii* were believed to have originated from cat feces, which survived or bypassed sewage treatment and traveled to coastal waters through river systems.

**FETAL INFECTION ROUTES**

Many viruses and microbes are capable of crossing the placenta. For example, the TT circovirus and several polyomaviruses (BK, JC, and SV-40) are able to cross the placenta and infect the fetus, but do not cause critical fetal abnormalities (37). In the laboratory, influenza A virus has been detected in mouse pups born of mothers exposed to influenza A virus (38). Similarly, influenza RNA has been shown to persist for at least 3 months following birth of offspring to mice infected with influenza A/WSN/33 virus during pregnancy (39, 40). A fetus may become infected with varicella (chickenpox) from its mother in the early weeks of pregnancy, manifesting as congenital defects particularly involving the cerebellum (41–43). Human transplacental transmission is well demonstrated by cases of fetal infection with rubella virus (44), West Nile virus (45), toxoplasmosis (46), and bovine leukemia virus (47).

**NON-SPECIES-SPECIFIC INFECTION**

Mice, rats, and other rodents potentially carry non-species-specific infectious diseases (31, 48, 49). Rodents also harbor insects and arachnids such as fleas, mites, and lice that can facilitate transmission of viruses, bacteria, and parasites to and between humans (31, 49). Certain arthropods in the mite family are known to transmit blood-borne diseases like “*Rickettsia typhi*” from mice and rats to humans (49). Although mites typically are host specific, they will cross from one species to another if their choice of host is unavailable (50, 51). Fleas are capable of passing diseases by blood from rodents to humans including bubonic plague, typhus, bartonellosis, and tapeworm (31, 48). Exposure to the respiratory or urinary aerosol of rodents also poses risk for human infections (e.g., *Hantavirus*, *Mycoplasma pulmonis*, *Streptococcus pneumoniae*, *Staphylococcus aureus*, and *Pasteurella multocida*) (52, 53).

The ease with which rodents transmit infection to humans likely reflects the high degree of homology between the two species. Approximately 80% of mouse genes have a single identifiable ortholog in the human genome, while <1% do not have any detectable human homolog (54). Occupations such as farming and agricultural work that place workers and their families in direct contact with rodents and their excretions/secretions increase the risk for infection (53, 55). KIPvV and WUPyV polyomaviruses have been detected in samples from children with lower respiratory tract disease, however it remains unknown if these viruses jumped species from rodents to humans (56).

**INDIRECT AND OTHER INTERMEDIATE MODES OF TRANSMISSION**

Domestic farm animals and cats that feed on infected rodents or share habitat with reservoir species may indirectly expose humans to rodent-borne pathogens (31, 53). Antibody to the rodent *Hantavirus* has been found in cats (53, 57–60). A case-referent study conducted in China found a positive association between cat ownership and *Hantavirus* infection (61). In the case of orthopox infection, cats have been confirmed as an active transmitter from rodents to man (57, 59, 62–66). A prevalence of
4% was observed in an antibody study of orthopoxvirus infection in 200 free-roaming cats, most of which appeared to occur through bites sustained during hunting of rodents (59). Compared with other times of the year, a markedly increased incidence of feline poxvirus infection has been reported to occur during autumn, when the populations of many small mammals are at their largest and most active (67–69). Human infection from farm animals and cats/dogs typically results from direct contact with infected tissues, contaminated mucous membranes, and skin wounds; inhalation of respiratory secretions; and injuries sustained from infected animals (31).

Hantaviruses have been isolated from lung tissues of bats raising the possibility that these creatures also may pose a natural reservoir for the passing of rodent viruses to farm animals (53, 70). Bats appear to play a role as well in the infection of pigs with Menangle virus (71) and horses with Equine morbillivirus (72, 73). Notably, these mammals host a significantly greater number of zoonotic viruses per species than rodents (74). Bats also have been observed to have a greater number of interspecific “host switches” of paramyxoviruses to other mammals, compared with rodents, birds, primates, carnivores, and cetartiodactyla (74). Among humans, bats have been associated with the spread of severe acute respiratory syndrome-like corona viruses, Ebola and Marburg filoviruses, and Hendra and Nipah paramyxoviruses (74). Because of habitat encroachment and the proclivity of bats to roost in pitched roof spaces of buildings, humans are increasingly at risk for exposure to bat excreta (74).

Other intermediate modes for the transmission of infectious agents include insects and birds. Flying insects, for example, may feed on animal manure and later contaminate nearby food and water sources intended for human consumption, especially in the agricultural setting (48). Mosquitoes are known to transmit West Nile virus and Japanese B encephalitis from avian and horse hosts to humans (75–77) and conceivably may act in an analogous fashion as a carrier of oncogenic viruses between animals and humans. Lice, ticks, and other bloodsucking insects also can serve as mobile vectors to transfer neurotropic viruses (e.g., Alphavirus and Flavivirus) from reservoir host animals to other susceptible species including humans (31, 78). Wild ducks and migratory birds shed virus in their feces as they fly south in the fall, contaminating water on farms where chickens, turkeys, and pigs are raised (79).

LOW OR NEGATIVE ANTIBODY RESPONSE
Viruses can induce cancer in species outside their natural hosts (80–82). However, not all viruses, especially those of avian origin, easily transmit directly to humans in nature (79). Canadian wildlife personnel, who handle wild ducks when they are shedding high levels of virus, consistently have negative viral and serologic assays (83). Volunteers inoculated with high doses of avian subtypes H4N8, H6N1, and H10N7 failed to produce a detectable antibody response, even though ~25% shed virus and had mild clinical symptoms (83). Only 2 of 27 persons in a sample of chicken farm laborers and residents regularly in contact with infected chickens and wildfowl tested positive for antibodies to Rous sarcoma virus-Bryan (RSV-B) (84). On the other hand, pigs have receptors for both avian and human viral strains and likely serve as a “mixing vessel” for genetic reassortment and subsequent introduction of genes into humans (85, 86).

DIRECT HUMAN TRANSMISSION
Avian viruses can directly infect humans without passing through pigs as evidenced by the recent outbreaks of influenza A strains H5N1 (87–88) and H9N2 (89) in Asia, and H7N7 (90) in Netherlands (78, 91). The history of human infection closely parallels the domestication of the duck, which introduced respiratory-type influenza A viruses into the “farmyard (91).” H5 subtype infection initially was detected in human sera from rural China, a region known for poultry farming (67, 92). Approximately 30% of poultry workers sampled around the time of the 1999 H9N2 flu outbreak in Hong Kong were found to be seropositive for antibodies to the virus (89). H9N2 viruses are prevalent in domestic poultry such as chickens, ducks, geese, quail, and pigeons throughout Asia (89). In Netherlands outbreak, investigators reported a high infection rate of H7N7 among people directly involved in handling infected poultry and noted evidence for person-to-person transmission (90).

ANIMAL INFECTION
Influenza viruses have been found to occur throughout the mammalian kingdom in nature (92, 93) and are capable of infecting (either naturally or experimentally) animals commonly found on farms including cattle, sheep, goats, horses, mice, and cats (53, 93–95). In Vietnam, civets (a cat-like species) have tested positive for H5N1 virus (96). Inoculation with various Influenza A strains (e.g., H5N1/97, H9N2, H5N3, A/tern/So, Africa/61, and A/turkey/England/63) can cause infection of mice (93, 97–100) and cross the blood–brain barrier (97–99, 101). The naturally occurring equine H7N7 influenza virus is highly pathogenic in laboratory mice, requiring no adaptation (99, 102).

SPECIES JUMPING
The transmission route for over 200 (~14%) human pathogens remains unknown and zoonotic transfer or species jumping cannot be ruled out as a potential source of infection (1). Among known routes, infected feces and urine remain one of the most prominent modes of transmission whether by primary or secondary sources. Animals that are not susceptible to infection may still spread disease to other animals and humans if they become contaminated and share water or food sources (103). However, for domestic poultry, secondary transmission often is associated with human involvement in the agriculture setting through personnel and/or animal movement, food delivery, and the use of fomites (103). Because poultry often do not display clinical signs of infection they may not be quarantined from other animals or their human caretakers (103). Furthermore, the increasing global demand for poultry and meat products coupled with improvements in transportation, have propagated closer contact between humans and farm animals (104).

EPIDEMIOLOGY
LACK OF SPECIFIC CANCER RISK ACROSS STUDIES
Epidemiologic evidence in support of an animal-transmitted, viral or bacterial etiology for human cancers is limited and often inconsistent with respect to specific cancer risk. For example, exposure
to farm animals or manure has been associated in some studies with childhood (105, 106) but not adult brain tumors (107, 108). A prospective cohort of 20,132 poultry, slaughterhouse workers (a group with frequent exposure to avian leucosis/sarcoma, reticuloendotheliosis, and Marek’s disease viruses) were observed to have a statistically significant excess of several cancers including lung [standardized mortality ratio (SMR) = 1.6, 95% CI = 1.3–1.7], cervix (SMR = 2.2, 95% CI = 1.3–3.5), penis (SMR = 8.6, 95% CI = 1.0–31.1), brain/nervous system (SMR = 1.7, 95% CI = 1.1–2.4), lymphoid leukemia (SMR = 2.2, 95% CI = 1.1–4.1), and monocytic leukemia (SMR = 9.2, 95% CI = 1.1–33.4) (7). Results of the Agricultural Health Study, a prospective cohort study of 49,884 male farmers, observed a statistically significant relative risk (RR) for non-Hodgkin lymphoma (RR = 1.6, 95% CI = 1.0–2.4) among farmers who raised poultry (109). However, in contrast to the above study of poultry, slaughterhouse workers, the RR among farmers who raised poultry was not significantly increased for either lung cancer or leukemia. Performing veterinary services also was associated with a statistically significant excess of Hodgkin lymphoma (RR = 12.2, 95% CI = 1.6–96.3) in the Agricultural Health Study. However, an increased risk for lymphoma among veterinarians has not been observed in other studies (110, 111).

**VIRAL EXPOSURE**

In a medical record-based study of 83 cases and 166 referents individually matched on date of birth, sex, and hospital of birth, children of mothers who had documented evidence of a clinically diagnosed viral infection during pregnancy had an 11-fold odds ratio (OR) [confidence interval (CI) = 1.1–503.2] for childhood neoplasm compared with unexposed mothers (112). However, none of the noted viral infections (mumps, varicella, herpes zoster, and rubella) were related to animal exposure. Similarly, a 2.4-fold OR (CI = 1.5–4.0, 25 cases) for childhood brain tumors (CBT) was observed in a nested (within Swedish birth-cohorts 1973–1989) case-referent (545 cases, 2798 referents) study of children born to mothers who reported a wide variety of neonatal viral and bacterial infections during the pregnancy of the index child (113). Significantly increased risk estimates were specifically observed for CBT subtypes “low-grade astrocytoma” (OR = 2.7, CI = 1.2–5.8) and “high-grade astrocytoma” (OR = 5.0, CI = 1.0–24.8). Neonatal urinary tract infections were associated with a 7.5-fold OR (CI = 1.3–44.9) for low-grade astrocytoma. This is in contrast to other case-referent studies examining vaginal and genitourinary infections during pregnancy, which did not observe a statistically significant increased OR for CBT (106, 112, 114).

**BACTERIAL EXPOSURE**

Cancer also has been linked with exposure to animal bacteria. In an ecologic, medical geographic study, a statistically significant 1.8-fold increase in brain cancer risk was observed for countries with increasing prevalence (4–67%) of *T. gondii* infection (115). In contrast, a case-referent study of brain cancer conducted in Australia observed no difference between participants with glioma (n = 117) and referents (n = 415) in the prevalence of antibody test-positivity (35% test-positive in glioma versus 33% in referents; age-, sex-, and center-adjusted OR = 1.00, 95% CI: 0.64–1.56) (33). While it is difficult to compare the results of the above two studies given their different study designs, ecologic studies generally are considered to provide the weakest form of epidemiologic evidence (116). For example, the ecologic link between brain cancer and *T. gondii* infection is difficult to interpret given the lack of patient-level information on histologic subtype and pathologic grade. Similarly, ecologic findings of increased cancer risk among dairy farmers must be carefully weighed against the absence of antibodies to bovine leukemia virus in these populations (117).

**RANDOM CANCER SUSCEPTIBILITY**

While few epidemiologic studies alone provide convincing evidence of an animal microbial basis for cancer, the data nevertheless are suggestive of a possible effect when examined as a whole. Perhaps the excess risk observed for some cancers, but not necessarily the same cancers across studies, may reflect a random cancer susceptibility to infection and inflammation rather than a specific microbe-cancer relationship. By analogy, animal viruses and bacteria may represent a chambermaid’s master key, capable of opening all hotel doors, but only if left unlocked by the guests.

**EXPLANATION OF INCONSISTENT FINDINGS**

Residual confounding or the lack of adjustment for factors such as population mixing, seasonality, climate, pesticides, medications, diet, and genetics may explain some of the conflicting and inconsistent results observed in the literature. Additionally, statistical results often are not appropriately adjusted for multiple comparisons and post hoc subset analyses, as was the case in a study reporting a statistically significant association between *T. gondii* infection and meningioma risk (33). Gene-environment interaction and the interplay between genes also are important puzzle pieces frequently missing from epidemiology studies of complex diseases such as cancer.

In the case of a “Hit and Run” virus, where the exposure theoretically occurred as an isolated event many years in the past, recall bias may have hindered the ability to establish causality or a temporal connection between a viral or bacterial exposure and cancer. A lack of study power, imprecision of point estimates, misclassification error, reverse causality, or selection bias represent other factors that may explain the inconsistencies observed across some studies and should be carefully considered with evaluating results (109, 112). Furthermore, the use of potentially carcinogenic disinfectant agents or cleaning compounds and must be taken into account as related etiologic exposures that might partially or fully explain a positive association between animal related microbial exposures and cancer.

**CONCLUSION**

While the general population is commonly exposed to animal viruses and bacteria, many of which are known to cause cancer in animals, the etiologic role of these exposures in human cancer remains speculative. For example, animal oncoviruses generally are species specific and do not infect or replicate easily in humans. Nevertheless, animal viruses conceivably may cause cancer in humans analogous to human and simian polyomaviruses causing tumors in non-permissive rodents. Epidemiologic studies to date have provided little evidence that animal viruses and bacteria cause
human cancer. Future studies will need to address the complex nature of cancer taking into account multiple interacting risk factors, and perhaps a non-stationary stochastic risk that contradicts conventional research design. The latter may be especially true given the waxing and waning behavior of viruses and bacteria. The same infectious agent may present and react differently depending on a host of factors including geography, seasona variation and climate, population density, and herd immunity. Travel, hygiene, and cultural variation in food consumption and preparation among individuals further complicate the epidemiologic study in this field.

AUTHOR CONTRIBUTIONS

Jimmy T. Efird: conception of manuscript, drafting of manuscript, critical revision of manuscript, and final approval of manuscript. Stephen W. Davies: critical revision of manuscript and final approval of manuscript. Wesley T. O’Neal: critical revision of manuscript and final approval of manuscript. Ethan J. Anderson, critical revision of manuscript and final approval of manuscript.

ACKNOWLEDGMENTS

Katherine T. Jones offered valuable comments during the writing of this manuscript and her contributions are greatly appreciated. This work was supported in part by NIH grant 2 G12 RR003061-2.

REFERENCES

1. Taylor LH, Latham SM, Woolhouse ME. Risk factors for human disease emergence. Philos Trans R Soc Lond B Biol Sci (2001) 356(1411):583–9. doi:10.1098/rstb.2001.0888
2. Parsonnet J. Introduction. In: Parsonnet J, editor. Microbes and Malignancy: Infection as a Cause of Human Cancers. New York: Oxford University Press (1999), p. 3–15.
3. Feng M, Zhao J, Petek H. Atomlike, hollow-core-bound molecular orbitals of C60. Science (2000) 320(5874):359–62. doi:10.1126/science.115866
4. Parkin DM. The global health burden of infectious-associated cancers in the year 2002. Int J Cancer (2006) 118(12):3030–44. doi:10.1002/ijc.21731
5. Tustin R. Ovine jaagsiekte. J S Afr Vet Med Assoc (1969) 40:3–23.
6. Weiss R. Viruses and cancer. In: Nesse R, editor. Infection as a Cause of Human Cancers Taking into Account Multiple Interacting Risk Factors – and an Associated Parasitic Agent. Nature (1948) 161(4090):441. doi:10.1038/161441a0
7. Ratcliffe HL, Worth CB. Toxoplasmosis of captive wild birds and mammals. Am J Pathol (1951) 7(4):655–67.
8. Barlok J. Toxoplasmosis of sheep in the lowland region of Western Slovakia. Vet Med (Praha) (1977) 24(5):285–94.
9. Anderson LM, Diwan BA, Fear NT, Roman E. Critical windows of exposure for children's health: cancer in human epidemiological studies and neoplasms in experimental animal models. Environ Health Perspect (2000) 108(Suppl 3):573–94. doi:10.1289/ehp.00108s3573
10. Rice JM, Ward JM. Age dependence of susceptibility to carcinogenesis in the nervous system. Am J Acad Sci (1982) 381:274–89. doi:10.1111/j.1749-6632.1982.tb05392.x
11. Belman D. Chronic host-parasite interactions. In: Parsonnet J, editor. Microbes and Malignancy: Infection as a Cause of Human Cancers. New York: Oxford Press (1999), p. 19–34.
12. Rice JM, Rehm S, Donovan PJ, Perantonio AO. Comparative transplacental carcinogenicity by directly acting and metabolism-dependent alkylating agents in rodents and nonhuman primates. DARC Sci Publ (1989) 96:17–34.
13. Martin B. Zoonotic Diseases: Institutional Animal Care and Use Committee. Available from: http://research.ucsd.edu/connect/pmd/disease.html
14. Clark S, Rylander R, Larsson L. Airborne bacteria, endotoxin and fungi in dust in poultry and swine confinement buildings. Am Ind Hyg Assoc J (1983) 44(7):537–41. doi:10.1080/15298683819405265
15. Ryan P, Hurley SF, Johnson AM, Salzberg M, Lee MW, North JB, et al. Tumours of the feline thymus. Toxoplasma gondii. Int J Epidemiol (1993) 22(3):412–9. doi:10.1093/ije/22.3.412
16. Centers for Disease Control and Prevention. CDC FactSheet: Toxoplasmosis Infection. Division of Parasite Diseases. Centers for Disease Control and Prevention (2004). Available from: www.cdc.gov/ncidod/dpd/parasites/toxoplasmosis/2004_pdf_toxoplasmosis.pdf
17. Hooke M, Saif D. Toxoplasmosis. Available from: http://www.emedicine.com/med/topic294.htm
18. Jones JL, Dargelos V, Roberts J, Press C, Remington IS, Montoya JG. Risk factors for Toxoplasma gondii infection in the United States. Clin Infect Dis (2009) 49(6):878–84. doi:10.1086/605433
19. Shirmizu B, Yu Q, Hu N, Yanagihara R, Nerurkar VR. Investigation of TT virus in the etiology of pediatric acute lymphoblastic leukemia. Pediatr Hema tol Oncol (2002) 19(8):543–51. doi:10.1088/0888-0018/19(9)/7/039
20. Siem RA, Ly H, Imagawa DT, Adams JM. Influenza virus infections in pregnant mice. J Neurophil Exp Neurol (1960) 19:125–9. doi:10.1007/000572-19600100-00013
21. Aronsson F, Lannebo C, Pauar M, Brack J, Kristensson K, Karlsson H. Persistence of viral RNA in the brain of offspring to mice infected with
Efird et al.

Viruses, bacteria, and cancer

influenza A/WSN/33 virus during pregnancy. J Neurovirol (2002) 8(4):353–7.

doi:10.1080/13555282002910480

Baldwin RT, Preston-Martin S. Epidemiology of brain tumors in childhood – a review. Toxicol Appl Pharmacol (2004) 199(2):118–31. doi:10.1016/j.taap.2003.12.029

Lafortet EV, Lynch CL Jr. Multiple congenital defects following maternal varicella; report of a case. N Engl J Med (1947) 236(15):534–7. doi:10.1056/NEJM194704102361504

Rivirk R. Congenital varicella encephalomyelitis in surviving newborn. Am J Dis Child (1969) 117(2):231–5.

Juel-Jensen BE, MacCallum F. Herpes Simplex Varicella and Zoster: Clinical Manifestations and Treatment. Philadelphia: Lipincott (1972).

Lee JY, Bowden DS. Rubella virus replication and links to teratogenicity. Clin Microbiol Rev (2000) 13(4):571–87. doi:10.1128/CMR.13.4.571-587.2000

Centers for Disease Control and Prevention (CDC). Laboratory-acquired West Nile virus infections – United States, 2002. MMWR Morb Mortal Wkly Rep (2002) 51(50):1133–5. Available from: http://www.cdc.gov/mmwr/preview/mmwrhtml/mm5150a2.htm

Lopez A, Dietz VJ, Wilson M, Navin TR, Jones JL. Preventing congenital toxoplasmosis. MMWR Recomm Rep. (2000) 49(RR):2:56–8. Available from: http://www.cdc.gov/cpr/mmwr/preview/mmwrhtml/rr00025s.htm

Van der Maaten MJ, Miller JM, Schmerr MJ. In utero transmission of bovine leukemia virus. Am J Vet Res (1981) 42(6):1052–4.

Williams BH. Zoonoses of small mammals. Clinician’s Brief (2013). 53–6.

McArthur J. Diseases of rats. Clinician’s Brief (2013). 52

Simmons JH, Riley LK. Hantaviruses: an overview. Clinician’s Brief (2013).

Waterston RH, Lindblad-Tow K, Birney E, Rogers J, Abril JF, Agarwal P, et al. Initial sequencing and comparative analysis of the mouse genome. Nature (2002) 420(6915):520–62. doi:10.1038/nature01262

Ogg B. Mites: medically important to humans. Zoonotic Disease Tutorial. Department of Pathological Sciences, School of Veterinary Medicine, University of Wisconsin-Madison (2004). Available from: http://www.vetmed.wisc.edu/pbs/zoonoses/

McArthur J. What’s your buggins your rats and mice? Rat and Mouse Gazette (1999). Available from: http://home.towson.edu/att/rattlvr/bugging1.htm

Waterston RH, Lindblad-Toh K, Birney E, Rogers J, Abril JF, Agarwal P, et al. Initial sequencing and comparative analysis of the mouse genome. Nature (2002) 420(6915):520–62. doi:10.1038/nature01262

Wright J. Diseases of rats. Clinician’s Brief (2013). 52

Pether JV, Trevains PH, Harrison SR, Baxby D, Bennett M, Gibb AP. Cowpox infection from cat to man. Lancet (1986) 1(8471):38–9. doi:10.1016/S0140-6736(86)91966-1

Bennett M, Gaskell CJ, Gaskell RM, Baxby D, Gruffydd-Jones TJ. Fxovirus infection in the domestic cat: some clinical and epidemiological observations. Vet Rec (1986) 118(4):387–90. doi:10.1136/vet.r.118.14.387

Chitty D, Phipps E. Seasonal changes in survival in mixed populations of two species of vole. J Anim Ecol (1986) 55:313–31. doi:10.2307/2398

Gibson D, Delany MJ. The population ecology of small rodents in Pen- nine woodlands. J Zool (1984) 203:63–85. doi:10.1111/j.1469-7998.1984.tb00644.x

Kim GR, Lee YT, Park CH. A new natural reservoir of Hantavirus: isolation of Hantaviruses from lungs of bats. Arch Virol (1994) 134(1):85–95. doi:10.1007/BF01379109

Philby AW, Kirkland JD, Ross AD, Davis RJ, Gleeson AB, Love RJ, et al. An apparently new virus (family Paramyxoviridae) infectious for pigs, humans, and fruit bats. Emerg Infect Dis (1998) 4(2):269–71. doi:10.3201/eid0402.980214

Young PL, Halpin K, Sellick PW, Field H, Gravel JL, Kelly MA, et al. Sero-logic evidence for the presence in Peromyscus of a paramyxovirus related to equine morbillivirus. Emerg Infect Dis (1996) 2(3):239–40. doi:10.3201/ eid0203.960315

Young P. Possible reservoir host of equine morbillivirus identified. Commun Dis Intell (1996) 20:262.

Luis AD, Hayman DT, O’Shea TJ, Cryan PM, Gilbert AT, Pulliam JR, et al. A comparison of bats and rodents as reservoirs of zoonotic viruses: are bats special? Proc Biol Sci (2013) 280(156):20122753. doi:10.1098/rspb.2012.2753.

Kumar N. West Nile virus: epidemiology and ecology in North America. Adv Virus Res (2003) 61:185–234. doi:10.1016/S0065-3527(03)60005-5

Petersen LR, Marfin AA, Gubler DJ. West Nile virus. JAMA (2003) 290(4):524–8. doi:10.1001/jama.290.4.524

Campbell GL, Marlin AA, Lanciotti RS, Gubler DJ, West Nile virus. Lancet Infect Dis (2002) 2(9):519–29. doi:10.1016/S1473-3099(02)00368-7

Olsen C. Zoonotic Disease Tutorial. Department of Pathological Sciences, School of Veterinary Medicine, University of Wisconsin-Madison (2004). Available from: http://www.vetmed.wisc.edu/pbs/zoonoses/

Webster RG, Shortridge KF, Kawaoka Y. Influenza: interspecies transmission and emergence of new pandemics. EMS Immunol Med Microbiol (1997) 18(4):275–9. doi:10.1111/j.1574-695X.1997.tb01056.x

Hoss HE, Olson C. Infectivity of bovine C-type (leukemia) virus for sheep and goats. Am J Vet Res (1974) 35(5):633–7.

McClure HM, Keeling ME, Custer RP, Marshak RR, Abt DA, Ferrer JF. Erythroblastosis in two infant chimpanzees fed milk from cows naturally infected with the bovine C-type virus. Cancer Res (1974) 34(10):2745–75.

Van Der Maaten MJ, Miller JM. Induction of lymphoid tumors in sheep with cell-free preparations of bovine leukemia virus. Bibl Haematol (1975) 43:377–9.

Bearse AW, Webster RG. Replication of avian influenza viruses in humans. Arch Virol (1991) 119(1):57–42. doi:10.1007/BF01341421

Morgan HR. Antibodies for Rous sarcoma virus (Bryan) in fowl, animal, and human populations of East Africa. II. Antibodies in domestic chickens, wild-fowl, pimates, and man in Kenya, and antibodies for Burkitt lymphoma cells in man. J Natl Cancer Inst (1967) 39(6):1229–34.

Kida H, Ito T, Yasuda J, Shimizu Y, Itakura C, Shortridge KF, et al. Evidence for human-to-human transmission of avian influenza A (H9N2) virus from poultry to humans. J Infect Dis (1985) 152(9):1727–32. doi:10.1093/infdis/152.9.1727

Subbarao K, Klimov A, Katz J, Regnery H, Lim W, Hall H, et al. Characteriza-

tion of an avian influenza A (H5N1) virus isolated from a child with a fatal respiratory illness. Science (1998) 279(5349):393–6. doi:10.1126/science.279.5349.393

Uyeki TM, Chong YH, Katz JM, Lim W, Ho YY, Wang SS, et al. Lack of evidence for human-to-human transmission of avian influenza A (H7N2)
Viruses, bacteria, and cancer

children: a report from the Children's Cancer Group. Cancer Epidemiol Biomarkers Prev (1994) 3(3):197–204.
97. Gemberge F, Little J, Golonca M, Azkan A, Preston-Martin S, Schlehofer B, et al. Contacts with animals and humans as risk factors for adult brain tumours. An international case-control study. Eur J Cancer (2002) 38(5):696–704. doi:10.1016/S0959-8049(01)00423–3
98. Ruder AM, Carreon T, Butler MA, Calvert GM, Davis-King KE, Waters MA, et al. Exposure to farm crops, livestock, and farm tasks and risk of glioma: the Upper Midwest Health Study. Am J Epidemiol (2009) 169(12):1479–91. doi:10.1093/aje/kwp075
99. Beane Freeman LE, Deroos AJ, Koutros S, Blair A, Ward MH, Alavanja M, et al. Poultry and livestock exposure and cancer risk among farmers in the agricultural health study. Cancer Causes Control (2012) 23(5):663–70. doi:10.1007/s10552-012-9921-1
100. Fasal E, Jackson FW, Klauber MR. Mortality in California veterinarians. J Chronic Dis (1966) 19(3):293–306. doi:10.1016/0021-9681(66)90133-0
101. Kinlen LJ. Mortality among British veterinary surgeons. Br Med J (Clin Res Ed) (1983) 287(6398):1017–9. doi:10.1136/bmj.287.6398.1017
102. Peer NT, Roman E, Ansell F, Bull D. Malignant neoplasms of the brain during childhood: the role of prenatal and neonatal factors (United Kingdom). Cancer Causes Control (2003) 14(2):443–9. doi:10.1023/A:1012102542589
103. Linet MS, Gridley G, Cnattingius S, Nicholson HS, Martinsson U, Glimelius B, et al. Maternal and perinatal risk factors for childhood brain tumors (Sweden). Cancer Causes Control (1996) 7(4):437–48. doi:10.1007/BF00052670
104. McKinney PA, Juszczak E, Findlay E, Smith K, Thomson CS. Pre- and perinatal risk factors for childhood leukaemia and other malignancies: a Scottish case control study. Br J Cancer (1999) 80(11):1844–51. doi:10.1038/sj.bjc.66906009
105. Thomas F, Lafferty KD, Brodeur J, Elguero E, Gauthier-Clerc M, Misse D. Incidence of adult brain cancers is higher in countries where the protozoan parasite Toxoplasma gondii is common. Biol Lett (2012) 8(1):101–3. doi:10.1098/rsbl.2011.0588
106. Greenland S, Morgenstern H. Ecological bias, confounding, and effect modification. Int J Epidemiol (1989) 18(1):269–74. doi:10.1093/ije/18.1.269
107. Blair A, Malker H, Cantor KP, Burmester L, Wiklund K. Cancer among farmers. A review. Scand J Work Environ Health (1985) 11(6):397–407. doi:10.5271/sweh.2208

Conflict of Interest Statement: The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

Received: 11 September 2013; accepted: 31 January 2014; published online: 13 February 2014.

Citation: Efird JT, Davies SW, O’Neal WT and Anderson EJ (2014) Animal viruses, bacteria, and cancer: a brief commentary. Front. Public Health. 2:4. doi:10.3389/fpubh.2014.00014 This article was submitted to Epidemiology, a section of the journal Frontiers in Public Health. Copyright © 2014 Efird, Davies, O’Neal and Anderson. This is an open-access article distributed under the terms of the Creative Commons Attribution License (CC BY). The use, distribution or reproduction in other forums is permitted, provided the original author(s) or licensor are credited and that the original publication in this journal is cited, in accordance with accepted academic practice. No use, distribution or reproduction is permitted which does not comply with these terms.