Widespread Outbreaks of a Subtle Condition Leading to Hospitalisation and Death

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Abstract

Long-term cycles in healthcare costs, medical admissions, hospital bed occupancy coupled with periods of increased deaths can be discerned in western health care systems for which there has never been an adequate explanation. A new type of immune-based infectious outbreak, operating via one of the many ubiquitous and persistent viruses appears to be occurring. Given the general difficulty of reaching a definitive diagnosis in the largely elderly population presenting to both primary and secondary care in western countries has led to these outbreaks remaining largely unrecognised. It is tentatively proposed that the ubiquitous herpes virus Cytomegalovirus (CMV) could be implicated in the observed long-term patterns of deaths, hospitalisation and wider costs.

Keywords: Immune impairment; Infectious outbreak; Cytomegalovirus; Aged populations; Death; Vague symptoms

Introduction

Since the 1960’s emergency admissions to hospital of a mainly medical nature in all western countries have increased in a manner far in excess of that expected from demographic changes in the population [1-4]. In order to explain this gap the various studies have implicated social and behavioural factors such as health care consumerism, more conservative General Practitioner (GP) and hospital consultant behaviour, the breakdown of the family, increasing numbers of elderly living alone and failings in the processes and organisation of health and social care – although in practice it would be exceedingly difficult to quantify the contribution from each of these factors [4]. However, no one seems to have posed the obvious question; could it be that there is a genuine trend to increasing poor health? Is it a co-incidence that illnesses such as diabetes, allergies, asthma and other immune syndromes have apparently increased in parallel with these unexpected increases in medical emergency admissions? [5-10].

Elderly Populations

In all western countries the generally aging population has resulted in the situation where the elderly account for the bulk of all primary and secondary care encounters and this is illustrated in Figure 1 for emergency hospital admission. It can be seen that 50% of adult acute hospital admissions are for those aged over 65 years and this proportion is increasing with time. This presents a unique set of problems regarding the correct diagnosis of the cause for the presenting symptoms. In the elderly, infection often manifests as changes in cognitive function (including depression) and physical function rather than the classical signs of infection such as fever seen in the young [11]. For example, in one study conducted in Nottingham, England during 2009 over 50% of patients aged over 70 years were cognitively impaired (27% with delirium, 32% depressed, 21% apathy, 17% delusions/agitation/aggression) [12]. Chronic inflammatory conditions associated with high levels of circulating cytokines lead to muscle wastage and disability (irrespective of age), although in the elderly the natural innate inflammatory responses are enhanced and prolonged [11,13]. Assessment scores for both cognitive and physical function are therefore good predictors of emergency department attendance [14] and of infection and/or inflammation and these measures often continue to decline after the immediately apparent
cause of the ‘infection’ (urinary tract infection, pneumonia, etc) has been treated [11].

In this context the presence of a subtle type of infectious immune impairment would be exceedingly difficult to discern.

An Immune-Based Syndrome

In 1981 the landmark paper by Michael Gottlieb, Joel Weisman and others described an association of unexpected infections in previously healthy young men which is now known as HIV/AIDS [15]. Are there a set of similar associations behind the seemingly inexorable rise in medical admissions seen over the past four decades? Using similar logic to that employed by Gottlieb et al a possible link has

![Figure 1: Age profile for adult acute emergency admissions.](image_url)
been explored in research which suggests that the increase in medical admissions (much like the cluster of unusual infections associated with HIV/AIDS) is characterised by all the features expected of an infectious outbreak rather than a more general trend which would arise if social and behavioural factors were the cause [4,16-20]:

1. The outbreaks occur at an interval of three to eight years with somewhere around five to six years being the most common.

2. At a local level (as opposed to a national level) these recurring increases occurs over a very short period (around 6 to 8 weeks) while at a national level the full spatial spread occurs over an 18 to 24 month period [21-25].

3. The onset of each ‘outbreak’ is characterised by a small but statistically significant increase in excess deaths. This immediate increase is smaller than the excess deaths which would typically characterise an influenza outbreak or similar however a cumulative increase in excess deaths over around one year can lead to total deaths in excess of a large influenza epidemic [22-25].

4. The increase is specific to aspects of medicine and mental health, i.e. admissions to the surgical specialties are largely unaffected.

5. Each outbreak leads to an approximate 10% step-like increase in medical admissions and bed occupancy with wider effects against total health care costs.

6. The increase is specific to a range of diagnoses which all have a common immune function linkage either via infection or inflammation but which are far wider in scope than those associated with HIV/AIDS [4]. This encompasses mental health issues which are dependent on immune function as an exacerbating factor and, as in AIDS, influence the trajectory of a small number of specific cancers [26].

7. Considerable regional [spatio-temporal] variation appears to be associated with each outbreak which is also suggestive of an infectious spread [4,22-25,27].

If this hypothesis is correct then wide spread disruption of health should occur and this should reflect in large scale changes in health care expenditures, especially given the generally higher costs associated with treatment of the elderly [28].

Health Care Costs

During the 1980’s the health insurance industry in the USA began to suspect that a curious cycle of profit and loss was occurring [29] and the evidence for this was formally reported in 1991 [30]. This cycle implied that the process of setting annual premiums via analysis of historical costs was subject to some form of sudden and unexplained change in the health of the population. Further research suggested that this cycle was linked to an approximate 6% [range 3% to 15%] increase in inflation-adjusted total costs [3,31] which will arise from a rapid change in the volume, case-mix and/or complexity of the health care contacts, the sum of which leads to the total cost of health insurance claims. While it is recognised that health insurance in the USA is typically more related to those of working age and generally the more affluent, the pattern of hospital admissions, occupied bed days and wider costs seen in other countries appears to mirror a cycle very similar, if not identical to, the health insurance underwriting cycle seen in the USA [3] as do total health care costs in the USA, i.e. including Medicare and Medicaid costs [32,33]. The last three of these potential infectious outbreaks occurred across the four countries of the UK (England, Scotland, Wales and Northern Ireland) toward the middle of 2002, 2007 and early in 2012 [4,23]. Figure 2 explores the impact on the cost of emergency admissions in England for the 2002 and 2007 outbreaks. As can be seen the rise in 2008/09 equivalent (i.e. inflation adjusted) costs was £627 million from the trough in 2001/02 to the peak in 2005/06 or £739 million from the trough in 2006/07 to the peak in

![Figure 2: Cost of emergency admissions in England.](image-url)
2009/10. Both of these figures are similar to a figure of £680 million estimated using the increase in occupied beds arising from the 2002 outbreak [27]. This sudden and huge increase in (largely) medical admission and bed occupancy has been proposed to be the main contributing factor to a cycle of surplus and deficit seen within the NHS over many years [32,33] and has curious similarities to the cycle of cost increases observed in the USA [34] and to a cycle in hospital admissions and bed demand seen in Canada and Australia [3,35]. The resulting inflation adjusted cost cycle seen in the USA is explored in Figure 3 where the gap from trough to peak ranges from 4% to 12% of total health care expenditure (average around 7%) and this equates to around $180 billion in 2008 equivalent costs. This estimate is close to a figure of 6% derived from reanalysis of the annual inflation-adjusted work of Born and Santerre [31] by this author [3]. The shoulder on the side of the first trough is likely to be associated with an outbreak occurring around 1974 or 1975 [3].

Clearly we are dealing with something having widespread and profound effects upon general health, including case mix and costs of health care [4].

Infectious Cycles

Outbreaks of all infectious diseases, where there is an element of acquired immunity, occur at a unique frequency particular to the infectious agent [34-42]. Certain viruses maintain a state of permanent or persistent infection in the host and this could account for the step-like change in emergency admission rates, which will arise from the pool of infected persons who could be experiencing some form of impaired or altered immunity.

Hence if the hypothesis of repeating international outbreaks is correct, then we must of necessity be looking for a ubiquitous virus known to establish a persistent infection but must additionally possess powerful immune modulating properties. In this respect, the ubiquitous herpes virus, Cytomegalovirus (CMV), could be implicated [4].

Cytomegalovirus

Up until recent years this virus has been largely regarded as only posing a risk to the developing foetus, where it is responsible for 40% of all congenital conditions [43,44] and the severely immune-compromised such as HIV/AIDS and transplant recipients [45,46]. Prevalence of this virus in Western countries increase with age with around 50% of the population CMV seropositive by age 15 to 34 and 80% by age 55 to 59 [44,47]. The multiple modes by which CMV could pose a serious problem to the seemingly immune competent has been recently reviewed along with the mechanisms by which CMV outbreaks could occur [4], however, the key point is that this virus possesses a powerful array of immune evasive and suppressing strategies against both adaptive and innate immune functions [45,48-51] leading to immuno suppression, chronic inflammation and autoimmunity [52-55] which will collectively be implicated in the observed higher all-cause mortality (hazard ratio of 1.2) for persons who are CMV seropositive [56]. The specific issues relating to mortality are discussed later. Due to the natural processes of immunoscenescence CMV has been particularly shown to be a risk factor in the elderly [57-60] where it contributes to what is known as the ‘Immune Risk Profile’ (IRP). Studies have shown that the healthy elderly are either not infected with CMV or have a significantly lower pro inflammatory state indicating that CMV is under strict immunological control [61-63].

Hence a general erosion of immune function and competence arising from life-long exposure to CMV is highly likely to lead to a host **Figure 3: Cycle in total health care expenditure in the USA.**
of non-specific symptoms which will be difficult to specifically attribute to CMV.

**Diagnostic Uncertainty**

In this respect, several studies have noted that a specific and unexplained increase in diagnoses which are described in Chapter R (signs and symptoms) of the International Classification of Disease (ICD) are associated with the increase in medical emergency admissions for the elderly [64-67] especially R00-09 (circulatory and respiratory) and R50-68 (general signs and symptoms) [68]. Re-analysis of the same data has demonstrated that particular signs and symptoms exhibit a characteristic step-like increase immediately after the 2002 and 2007 ‘outbreaks’ seen in the UK [4] and a similar ‘outbreak’ seen in Australia [69]. In the USA, a specific increase in emergency department attendances for ‘other and undefined diagnoses’ has been observed since 1993 and this translated into peaks in admission via the emergency department around 1993, 1997 and 2003. The highest increase was noted in those aged over 65, African-descent and requiring three or more medications [70]. Hence the general difficulty in assigning a definitive diagnosis in the elderly is consistent with the specific step increase in admissions for signs and symptoms and this may have masked the true underlying aetiology.

In general practice it is known that less than 20% of the most frequent diagnoses account for 80% of consultations, hence, just 20% of consultations relate to the 80% of less frequent and hence less familiar diagnoses. In general, 50% of diagnoses remain a description of symptoms, 40% are named syndromes and only 10% are a confirmed diagnosis [71]. This diagnostic ambiguity will almost certainly also apply in the acute context.

In addition to the more general ‘signs and symptoms’, there are an additional set of more specific diagnoses that appear to be implicated in a series of step-like increases in emergency admission and in the widest sense all such diagnoses have a common link with infection or inflammation, i.e. inflammatory immune responses and the consequences of these [4]. The point of relevance is that a combination of diagnostic ambiguity and the generally elderly age of patients would make it exceedingly difficult to detect a general immune-based syndrome.

For example, an elderly patient presenting with pneumonia would be diagnosed as one of the multiple forms of ‘pneumonia’ although the exact reason that the patient has pneumonia rather than remaining healthy may well be related to a CMV induced recent deterioration in the IRP or other measures of inflammation and immune competence and CMV re-activation in the lungs leading to a more specific diagnosis of CMV pneumonitis [72]. Is the possibility that urinary retention can arise from CMV infection widely appreciated? [73]. Would an elderly patient presenting with depression be prescribed anti-depressants if the doctors were aware that active CMV infection is associated with depression in the elderly [74] and that disturbances in immune function are now increasingly recognised as a primary cause of depression [75,76]. Would a link between cirrhosis of the liver and CMV be explored [77]? Indeed how many clinicians would be aware that CMV has been directly linked to breast cancer [78,79], Ross syndrome [80], chronic periodontitis [81], pre-eclampsia [82], inflammatory bowel disease [83]. These are but a small sample of a growing body of medical case studies involving CMV in a diverse range of more complex illnesses which are discussed further in a later section. Indeed a recent reviews of severe CMV infection in apparently immune competent patients has concluded that severe life-threatening complications may not be as rare as previously thought [84-86]. In critically ill patients CMV infection (without bacterial infection) gave relative risk of 4 (sepsis, respiratory failure, death), 5 (pneumonia, acute respiratory distress syndrome, multi organ failure), 6 (diarrhoea) and 11 (septic shock) [87]. The likelihood is that we are treating symptoms rather than root causes.

The following sections will investigate the common themes relating to the clinical aspects of CMV infections and how these may overlap with the observed infectious-like outbreaks.

**CMV Strains**

CMV displays genetic polymorphisms in multiple genes [88-90] with possible clinical significance. It has been recently proposed that this diverse range of polymorphisms may confer advantage in the ability to exploit a wide range of temporary through to permanent immune impairments present in all supposedly ‘immune competent’ persons [4].

CMV is able to rapidly mutate both in vivo and in vitro. Endothelial cell and leucocyte (polymorphonuclear & monocyte) tropism is shared by all clinical isolates but is missing in laboratory adapted strains [91] while long-term exposure to a range of anti-virals produces resistant strains [92].

The prevalence of different CMV strains appears to vary widely between countries. In Japan a mutation at codon 605 (D to E) is present in 92% of CMV infected children while this particular strain has a low prevalence in Western countries [93]. In India, Glycoprotein GB3 is the most frequent compared to GB1 in China and Hungary [94] while GB1 also occurred more frequently in a 1998 study in Iowa, USA [95]. While in Baltimore, USA in young women who had recently acquired CMV infection some 75% had a unique strain or a strain shared with only one other woman, 25% shared a common strain while 4% had multiple strains [96].

These different strains do have different properties in vivo and in vitro and the kinetics of cytolyis of CMV infected fibroblasts depends on the strain [97]. The ability of different CMV strains to produce active infection of endothelial cells (the most common clinical location for CMV infection and disease) depends on mutations in the UL133-UL138 locus of the CMV genome [98]. Strain specific interactions between CMV encoded proteins and epigenetic mechanisms (histone acetylation and deacetylation) have been observed [99] and posttraumatic stress disorder (PSTD) patients have specific epigenic modifications accompanied by higher levels of CMV antibodies [100].

Infection of infants showed clustering of disease types according to CMV Glycoprotein B genotypes [94] and another study has suggested that there is a 5-fold reduction in the risk of sequealae in congenitally infected infants for Glycoprotein GN-1 and GN-3a genotypes, however GN-4 genotypes appear to increase risk by 8-fold [101]. However in organ transplant recipients Immediate Early 1 (IE1) variants showed no discernable relationships [88]. In this respect Glycoprotein N appears to be more to do with CMV neutralisation by antibodies rather than more active CMV-manipulation of the immune system [102]. However in the latter study mixed strain infections demonstrated associations and outcomes that single-strain infections did not [88].

Multiple genotype infections are relatively common and up to six different genotypes have been detected in a single adult [89,90]. The role of different strains of CMV is important and infection of mice with multiple CMV strains leads to active infection of the salivary ducts (a potential source of salivary duct cancer) with the more pathogenic strains [103]. In lung transplant recipients single strain infections
are generally asymptomatic while mixed strain infections produce symptomatic infection [89].

Introduction of a new strain is a feasible reason for these outbreaks.

Mixed Infection

In just the same way that infection with multiple CMV strains leads to more clinically serious outcomes so also does multiple infection with other viruses, bacteria and fungi. For example, in the era prior to antivirals in bone marrow transplant patients with CMV pneumonia who exhibit lymphocytosis progress to recovery while those who had mixed bacterial or fungal infection with peripheral lymphopenia died [104]. In the critically ill patient those with a mixed CMV/bacterial infection (which occurred most often in the winter, with 4-times higher length of stay and peaked in 50 to 60 year olds) had very high risk ratios of 7 (death), 10 (diarrhoea, multi-organ failure) and 220 (septic shock) [87]. Patients with mixed Hepatitis C Virus (HCV) and CMV infection who do not spontaneously clear the HCV infection generally have higher levels of CMV IgG and IgM and have double the level of CMV DNA detection in serum [105]. A case of double encephalitis is described in a patient with dual infection with HSV and CMV [106]. Acute respiratory infections (RSV, rhinovirus, enterovirus) are statistically more frequent in children infected with CMV [107].

In critically ill patients reactivation of HSV then leads to a cascade of events leading to CMV re-activation [108] and presumably vice versa and of other herpesviruses such as Human Herpesvirus 6 (HHV-6) which can lead to further complications [109]. Such a cascade of intertwined effects is also observed in the ability of latent HHV-6, HSV and CMV infection to induce Chlamydiaid persistence via imbalanced oxidative stress [110]. Such cascades may form part of the proposed time-dependant shifts in GP referral for different dermatological conditions that appear to occur after these outbreaks [111].

It would appear that synergistic interactions with other resident infections are capable of enhancing the clinical consequences of a CMV-based outbreak.

Genetic Factors

As expected, genetic factors are also important. Those with a variety of schizophrenia (including neurophysiology) susceptibility genes have heightened susceptibility to CMV and several other pathogens [112]. Early life environment (epigenetic factors?) [99] has likewise been shown to be important in the susceptibility of individuals to acquire CMV infection [113]. There is evidence for the epigenetic regulation of cytomegalovirus gene expression [114] and CMV in turn modifies the epigenetic processes [115].

Hence both genetic and epigenetic factors will be involved in the differential sensitivity of individuals to a variety of CMV-mediated disease processes.

Gender

Men are generally known to be more susceptible to infection while women have a far greater risk of illness caused by an overactive immune system [116], hence at the same level of treatment immune-mediated chronic inflammatory diseases have a greater effect against women [117]. While such generalisations are useful it is of interest to note that there are distinct gender differences in the types of immune response determining susceptibility to CMV-mediated coronary artery disease with men mounting a far more inflammatory-based response centred around elevated levels of CRP while that in women is more subtle [118]. CMV antigenemia in cancer patients is higher in women [119] and women are known to have higher release of interferon and interleukin 2 in response to CMV infection [120]. In older women CMV antibody levels are associated with increased risk of diabetes, CVD, frailty and mortality (HR 2.8) [60].

The literature is conclusive that CMV seropositivity is higher in females [4,43,44,121] and it is a notable feature of the infectious outbreaks that emergency admission and death is higher in females [4,23]. Even more curiously each outbreak appears to initially elevate the gender ratio at birth (male: female) which is the opposite to what would normally be expected, i.e. the outbreak appears to have a particular effects which extends to the female foetuses [122]. CMV DNA is detected in 15% of >20 week stillborn and manifests as thrombotic vasculopathy in 60% of cases [123]. Congenital CMV mortality in the US is highly race specific with odds ratio of: Native American 2.3, African American 1.9, White American 1.0, Hispanic 0.96, Asian 0.5 [124]. Male/female differences are therefore likely and in congenital CMV infection females have double the risk of brain abnormalities [125]. Hence the observation of undulations in the gender ratio initiated by each outbreak is feasible.

Hence each outbreak is associated with generally higher admissions in women and a smaller and more transient increased loss of the female foetuses. Gender specific responses to CMV appear to be a likely explanation. Further research is needed to disentangle if this effect is specific to particular conditions as appears to be the case [126].

Death

It has been noted that each outbreak appears to be associated with a generalised increase in deaths which lasts for around 12 to 18 months following the onset of the outbreak [23-25]. This has been especially apparent during the most recent outbreak occurring in early 2012 in England, although slightly earlier in Scotland [24] with approximately 40,000 excess deaths since the onset of the outbreak [23]. Several large population studies have noted that those who have CMV infection show higher mortality than their non-CMV infected counterparts, especially in those with elevated CMV antibody levels who are also characterised by an exaggerated inflammatory response.

One study in elderly Latinos over a nine year period showed a fully-adjusted 35% increase in mortality due to Cardiovascular Disease (CVD) and a 19% increase in all-cause mortality for those in the highest quartile of CMV IgM titre. The effect appeared to be driven by inflammation with links to the levels of interleukin-6 (IL-6) and Tumor Necrosis Factor (TNF) [127].

Another study representative of the population of the USA over an 18 year period demonstrated that individuals who were CMV seropositive and with elevated levels of C-reactive protein (CRP) showed a 30% increase in all-cause and CVD-related mortality compared to CMV seropositive but low CRP subjects [56].

In the EPIC-Norfolk [England] study which ran from 1993 to 2011, i.e. the study spanned four outbreaks in 1993, 1996, 2002 and 2007, CMV infection was associated with a 16% increase in all-cause mortality (+6% cardiovascular disease, +13% cancer and +23% other causes of which 12% were respiratory, 16% gastrointestinal and 21% central nervous system leaving 60% across other body systems) [128]. While CMV infection alone was associated with a 16% increase in all-cause mortality it was noted that those with the highest levels of CMV-antibodies had a 26% increase in all-cause mortality during the study.
CMV infection of the heart has been known for some time to be associated with fatal myocarditis [129].

The fact that all-cause mortality reaches across a wide range of body systems and diagnoses in the above study is consistent with the observed increase in emergency admission for a variety of medical conditions [4] and increased deaths [25] which appear to be associated with these outbreaks. For example, in end stage renal disease (ESRD) CMV aggravates the contraction of CD4+ naïve T cells and increases the number of differentiated CD4+ and CD8+ memory cells [130,131]. In HIV disease CMV-specific CD8+ T cell responses were lower during recent HIV infection, higher during chronic untreated infection and higher still during long-term antiretroviral treatment [46]. It would appear that the ability of CMV to manipulate multiple innate and adaptive immune functions lies behind this increased all-cause mortality [45,48,51,53-55,132] and the emergence of diverse diseases including Alzheimer’s and type 2 diabetes in the elderly [133,134]. In fact there are now increasing reports of cases where CMV viremia has been the direct cause of death in a variety of clinical contexts [25,86]. Unsurprisingly the absence of CMV-mediated inflammatory responses are a characteristic of longevity in 85+ year olds [59,61-63].

How do we link the above to the approximate 3% increase in deaths which occur during the first one to two years of these outbreaks; CMV-induced inflammatory responses are obviously part of wider all-cause mortality and the introduction of a new strain of CMV (which may only lead to a small increase in the proportion of CMV seropositive) into the population (see previous section) would therefore lead to a much smaller increase in deaths. There is some evidence to suggest that the proportion of CMV seropositive individuals does indeed fluctuate over time [4] and the next step will be to ensure that future population studies take this possibility into account. Indeed the most recent 2012 outbreak in the UK has led to higher levels of increased death than in previous outbreaks and re-analysis of clinical samples collected before and during this period is probably a useful endeavour [4].

In England each outbreak leads approximately to an additional 420,000 admissions [4] and this would imply serious infection leading to hospitalisation in just 5% of adults over the age of 65, of which only 10% actually die. This figure would be achievable and is consistent with the generally erosive effects of CMV against health rather than a virus leading specifically to high mortality- which is not a desirable outcome from a viral infectious point of view.

Cancer

In recent years CMV has been increasingly implicated as both an oncomodulatory and oncogenic agent [52,78,135-141]. A recent study of the trends in new diagnoses of cancers between 1999 and 2007 in the UK has led to higher levels of increased death than in previous outbreaks and re-analysis of clinical samples collected before and during this period is probably a useful endeavour [4].

A review of diagnoses for hospital admissions where the patient eventually died in the period following the 2007 outbreak in England showed a high proportion of cancer patients [26]. In this respect low level of CMV infection in Glioblastomamultiforme (an aggressive brain tumour) is known to be associated with patient survival [143] and CMV viremia in cancer patients is known to vary with cancer location, type and ethnicity of the patient [119].

Hence the evidence points toward a recurring series of some form of infectious outbreak with effects against a range of cancers where CMV seems to be a common theme.

Cognitive Function

CMV is becoming increasingly recognised as a psychotrophic agent. This is unsurprising given the bi-directional relationships between depression and other illness behaviours and immune function [75,76]. In schizophrenia CMV seropositivity is known to be associated with visual search, working memory and psychomotor speed aspects of cognitive function while both CMV and HSV1 are associated with error making, a cognitive flexibility and executive function [144]. In healthy adults CMV appears to influence the degree of novelty seeking behaviour [145]. In the elderly active as opposed to dormant CMV infection has been associated with depression [74]. More recently CMV has been implicated as part of the link between stress, ageing and immunity [146]. In Alzheimers Disease (AD) CMV antibody levels are associated with extent of Neuro Fibrillary Tangles (NFT) and interferon-γ is only found in the Cerebrospinal Fluid (CSF) of CMV seropositive subjects [133].

Hence the observed increase in admissions relating to self-harm and other mental health issues during the proposed outbreaks is consistent with a role for CMV [4].

Inflammation

Numerous studies have demonstrated that it is the ability of CMV to provoke an inflammatory response in particular individuals rather than CMV infection per se that is the key to understanding why CMV leads to illness in some people more so than others. Various inflammatory markers are therefore elevated and it is this which could lead to the wide variety of illnesses observed in these outbreaks. Hence elevated CRP (in a dose dependant manner) is known to be associated with use of antidepressants and risk of hospitalization. Odds ratio 2.7 for antidepressant usage or 2.3 for hospitalization in those with CRP >10 mg/l of serum [147]. See above section.

Emergency admission to hospital for potentially preventable and other conditions is highest in the most deprived quintile (as is the proportion who are CMV seropositive [43,44,121]), those with one or more mental health conditions (previous section) and for those with four or more multi-morbidities [148]. CMV-mediated inflammation is almost certainly part of these risk factors.

Role of the Thymus

The thymus is central in immune homeostasis and generates self-tolerant naïve T cells as well as self-antigen specific natural T-regulatory cells. It is a unique site where the endocrine and immune systems interact [149]. As such thymus function is regulated by both somatotrope Growth Hormone (GH) and Insulin-like Growth Factor 1 (IGF-1) which is central in insulin specific auto reactive T-cell selection [150]. The production of thymulin, an anti-inflammatory hormone, also occurs in thymic epithelial cells. Production of this hormone is strongly influenced by the neuroendocrine system [151]. A recent review of the action of CMV in disease has implicated thymic function as a specific variable of interest [4]. Thymic output reaches a maximum around age one and thereafter declines with age in a generally exponential manner with a 10-fold reduction at around age 45 and a further 10-fold reduction by age 80 [152], however, for individuals of the same age thymic output varies by a factor of 10-fold [153] due to a variety of adverse environmental factors [154]. This age related decline
is enhanced by thymectomy [155], end stage renal disease [130,131] or in Severe Combined Immuno Deficiency (SCID) where bone marrow transplantation effectively restores thymic output [156]. In humans, ageing and loss of thymic function is often accompanied by impaired zinc status. In aged mice zinc supplementation has been shown to increase thymic output and reduces the expression of stem cell factor, a thymo- suppressive cytokine [157].

Given the role of CMV in provoking inflammation in certain individuals it is of interest to note that thymopoiesis in elderly humans has been correlated with a shift to neutrophilic and inflammatory status [158]. A more recent study by the same group has demonstrated that both thymic output and levels of CRP are independent predictors of all-cause mortality [159]. Clearance of CMV viremia following double umbilical cord blood transplantation for hematologic malignancies in adults has been shown to rely on T cell neogenesis via the thymus [160].

The epithelial cells of the thymus medulla secrete Macrophage-Derived Chemokines (MDC) that chemotactically attracts the immature thymocytes. These MDC are probably responsible for the maturation of the thymocytes during their migration from the cortex into the medulla.

Infection of the thymus with particular agents can therefore disrupt thymic function. Hence in mice infection with Francisella tularensis leads to severe thymic atrophy and CD4 and CD8 depletion [161]. Persistent infection of murine thymic epithelial cells with Coxsackie virus B4 decreases the production of IGF-2, a hormone involved in self-tolerance toward pancreatic islet β cells [162], while persistent infection of human thymic epithelial cells with the same virus increases epithelial cell proliferation and production of inflammatory cytokines [163]. It is of interest to note that in vitro infection of human thymic epithelial and monocyte cells by measles, while causing a transient increase in thymic output leads to a measure of epithelial apoptosis and a decrease in the size of the thymic cortex [164].

In line with the wider preference of CMV for epithelial cells [165] a number of studies have demonstrated specific effects of CMV against the epithelial layer in the thymus where an active and persistent infection produces distinct multinucleated cells [166]. Such infected cells show a significant reduction in the production of cells reactive with monoclonal antibodies specific for mesoderm-derived compounds and a reduction in IL-1 related antigen production [167].

Further to the theme of epithelial cells, individuals with inflammatory skin diseases typically show depressed thymic function. In particular those with atopic dermatitis had high variability in thymic output over time [168]. Unsurprisingly CMV is often associated with these conditions and increased GP referral to dermatology typically occurs during the proposed outbreaks [111].

Hence it appears highly likely that in those individuals where CMV has infected the thymus the resulting disruption of thymic functions could lead to the enhanced inflammatory responses which have been observed in some members of CMV seropositive populations.

**Sub-Acute Infection**

One of the big questions in CMV pathology is how the virus appears to exert its effects in the absence of a clinical level of infection. The first hint at an answer to this question comes in the observation that CMV is often associated with cancerous tissue [52,78,135-141], i.e. it is simply taking site-specific advantage of one of a variety of exploitable immune or physiological impairments [4].

Perhaps the biggest development in this area is the discovery that in patients with Inflammatory Bowel Disease (IBD) CMV can re-activate in the gut mucosa without causing detectable changes in serum viral (DNA) load or traditional antibody assays [169]. These ‘hidden’ re-activations could, however be detected using an assay for recently activated ‘effector’ CD8 T cells.

Hence site-specific CMV infections in the vasopressin producing (AVP) part of the anterior hypothalamus [170], anterior pituitary gland [171], pancreatic islets [172,173] and a host of other cell types and tissues can all induce a variety of diseases or modify the course of another disease without typical CMV clinical symptoms. See section on Acute Admission for further detail.

Evidence is therefore accumulating for a wide range of infection types from site-specific infections through to more clinically observable ‘typical’ infections.

**Acute Admission**

There is now a large body of case reports for patients admitted to hospital with a wide variety of symptoms where CMV is a causative or exacerbating factor. One review identified 207 case reports up to 2007 with gastrointestinal, respiratory and central nervous system infections being most common [86]. Table 1 [174-181] gives a limited selection of such case reports which illustrate some of the principles highlighted above. For those patients admitted to intensive care or burns units CMV is a well-recognised risk factor with high rates of re-activation or first time infection leading to serious viremia, other complications and even death [84,182-184]. Indeed CMV involvement in patients with sepsis or septic shock is highly likely [108,185,186] and should be investigated as a possibility.

Hence we can propose that typical CMV viremia (usually with fever, diarrhea, etc) is more to do with the balance of latency and re-activation in circulating fibroblasts and undifferentiated myeloid cells [187] while specific ‘hidden’ infection in a much wider range of cell types and tissues [188] regulates the expression and exacerbation of a wider range of conditions and diagnoses.

Given the ability of CMV to increase all-cause mortality across a very wide range of conditions and diagnoses and the wide range of conditions associated with increased hospital admission it would seem that CMV is therefore a good match with the observed increases in death and hospital admission (for a wide range of otherwise apparently unrelated conditions) observed in these outbreaks.

Having established a clinical basis for the proposed infectious outbreak the issue of spatio-temporal spread is crucial in demonstrating that a genuine infectious spread lies behind the phenomena.

**Spatio-Temporal Spread**

As with any persistent infection CMV is in a state of a continuous ‘epidemic’ as the virus is spread via contact with body fluids in a variety of ways [44,121]. As a consequence CMV mini-epidemics have been documented in households, neonatal intensive care units, geriatric units and renal transplantation units [189-192].

If we are dealing with the introduction of a new strain then the levels of CMV infected persons should change over time and a review of the available, but rather limited literature on this topic, suggests that this is indeed possible but requires further research [4]. However, study of the 2002, 2007 and 2012 outbreaks in the UK has demonstrated that there is a very definite spatial spread associated with each event.
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**Conclusions**

It would seem that we do indeed have a phenomena contributing to the unexplained rise in medical (and especially elderly) admissions and that this phenomena is capable to causing large undulations in health service contacts and total health care costs. CMV is likely to be implicated at multiple levels, as direct agent, indirect risk factor and via opportunistic immune assault, as witnessed in patients in the intensive care unit [183,184]. Evidence has been presented to show that CMV is capable of active (but generally subclinical) infection of a wide range of cell types and tissues and is more than capable of directly and indirectly causing the wide range of diagnoses observed to associated with increased health service contacts and deaths associated with the infectious outbreaks. Is it possible that the real need for the elderly is targeted anti-viral and/or immune restoring therapy to maintain optimum health and well-being [196] and/or CMV growth inhibitors such as Vitamin A, monolaurin and lactoferrin [197], rather than the reactive treatment of presenting symptoms which they currently receive? The role of the thymus requires far greater attention as does the differences in immune response between the two genders. Widespread disruption in western populations appears to be related to country- and/or racial-specific distribution of different CMV strains. It would seem that very large sums of expenditure and the alleviation of widespread poor health may rest on the answer to these questions.

**References**


