Schizophrenia Susceptibility Genes Directly Implicated in the Life Cycles of Pathogens: Cytomegalovirus, Influenza, Herpes simplex, Rubella, and Toxoplasma gondii

C.J. Carter¹,²
¹¹76 Downs Road, Hastings, East Sussex, TN34 2DZ, UK
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Many genes implicated in schizophrenia can be related to glutamatergic transmission and neuroplasticity, oligodendrocyte function, and other families clearly related to neurobiology and schizophrenia phenotypes. Others appear rather to be involved in the life cycles of the pathogens implicated in the disease. For example, aspartylglucosaminidase (AGA), PLA2, SIAT8B, GALNT7, or B3GAT1 metabolize chemical ligands to which the influenza virus, herpes simplex, cytomegalovirus (CMV), rubella, or Toxoplasma gondii bind. The epidermal growth factor receptor (EGR/EGFR) is used by the CMV to gain entry to cells, and a CMV gene codes for an interleukin (IL-10) mimic that binds the host cognate receptor, IL10R. The fibroblast growth factor receptor (FGFR1) is used by herpes simplex. KPNA3 and RANBP5 control the nuclear import of the influenza virus. Disrupted in schizophrenia 1 (DISC1) controls the microtubule network that is used by viruses as a route to the nucleus, while DTNBP1, MUTED, and BLOC1S3 regulate endosomal to lysosomal routing that is also important in viral traffic. Neuregulin 1 activates ERBB receptors releasing a factor, EBP1, known to inhibit the influenza virus transcriptase. Other viral or bacterial components bind to genes or proteins encoded by CALR, FEZ1, FYN, HSPA1B, IL2, HTR2A, KPNA3, MED12, MED15, MICB, NQO2, PAX6, PIK3C3, RANBP5, or TP53, while the cerebral infectivity of the herpes simplex virus is modified by Apolipoprotein E (APOE). Genes encoding for proteins related to the innate immune response, including cytokine related (CCR5, CSF2RA, CSF2RB, IL1B, IL1RN, IL2, IL3, IL3RA, IL4, IL10, IL10RA, IL18RAP, lymphotxin-alpha, tumor necrosis factor alpha [TNF]), human leukocyte antigen (HLA) antigens (HLA-A10, HLA-B, HLA-DRB1), and genes involved in antigen processing (angiotensin-converting enzyme and tripeptidyl peptidase 2) are all concerned with defense against invading pathogens.

Human microRNAs (Hsa-mir-198 and Hsa-mir-206) are predicted to bind to influenza, rubella, or poliovirus genes. Certain genes associated with schizophrenia, including those also concerned with neurophysiology, are intimately related to the life cycles of the pathogens implicated in the disease. Several genes may affect pathogen virulence, while the pathogens in turn may affect genes and processes relevant to the neurophysiology of schizophrenia. For such genes, the strength of association in genetic studies is likely to be conditioned by the presence of the pathogen, which varies in different populations at different times, a factor that may explain the heterogeneity that plagues such studies. This scenario also suggests that drugs or vaccines designed to eliminate the pathogens that so clearly interact with schizophrenia susceptibility genes could have a dramatic effect on the incidence of the disease.

Key words: influenza/herpes simplex/cytomegalovirus/rubella/toxoplasmosis/T. Gondii/pathogen/genes/gene environment interaction/host pathogen interaction/schizophrenia

Introduction

Over 240 gene variants have been implicated in schizophrenia in genetic association studies. Such studies are notoriously inconsistent, and problems of replication are the norm rather than the exception. It has, however, been noted that many of these genes can be assembled into discrete but interconnected signaling networks related to glutamatergic neurotransmission and neuronal plasticity, oligodendrocyte function, and oxidative stress, all relevant subpathologies or endophenotypes of schizophrenia. The more important genes, eg, neuregulin or DISC1,¹ appear to be those residing at key hubs of these signaling networks or those with multiple effects on several areas of the network.²

Several viral or bacterial pathogens as well as famine in prenatal and postnatal periods (see below) have also been implicated in schizophrenia. Such stressors activate an eif2alpha kinase signaling cascade, again composed of proteins encoded for by schizophrenia susceptibility candidates, that shuts off protein synthesis in response to stress. This is achieved by inhibition of the translation

¹To whom correspondence should be addressed; e-mail: chris_car@yahoo.com.

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initiation factor eif2b, whose malfunction has been associated with oligodendrocyte cell loss in vanishing white matter disease. Glutamatergic (eg, N-methyl-D-aspartic acid [NMDA] receptors) and growth factor signaling networks (eg, neuregulin 1 [NRG1], brain-derived neurotrophic factor [BDNF]) also control protein translation and synthesis via this network, and such effects are important in the control of synaptic plasticity and dendritic growth. These pathways, created by a lattice of susceptibility genes may represent key intrinsic signaling networks whose malfunction may contribute to many pathological features of schizophrenia including oligodendrocyte cell loss, reduced glutamatergic function, and impaired synaptic connectivity.

Several studies have shown that maternal viral or parasitic infection during pregnancy is an important risk factor for the subsequent development of schizophrenia in the adult offspring. The pathogens implicated in these maternal effects include influenza, herpes simplex virus (HSV)-2, rubella, and Toxoplasma gondii. A large Danish study has also reported that viral infection during early childhood (0–12 years), particularly with cytomegalovirus (CMV) or mumps are associated with an increased risk of psychosis. Seropositivity to HSV-1, related to cognitive deficits and cerebral gray matter changes has also been reported in adult schizophrenics and antibodies to HHV-6 or T. gondii prior to diagnosis in adults have also been associated with schizophrenia. Increased seropositivity for CMV has also been observed at presentation in first-episode schizophrenic patients. It would, thus, seem that prenatal, childhood, and adult infections could all act as risk factors, perhaps, in a synergistic or additive fashion. Further examples (borna virus, Borrelia burgdorferi, chlamydial infection, coxsackie B5, poliovirus, retroviruses) are listed at http://www.polygenicpathways.co.uk/. Herpes viruses (HSV-1 and HSV-2 and CMV) are neurotropic and known to be able to infect the central nervous system, although only HSV-1, HSV-2, CMV, and HHV-6 have been detected in schizophrenic brain tissue. The influenza virus is also capable of cerebral entry through the nasal route via olfactory nerves, at least in mice. Toxoplasma gondii is also capable of infecting the brain but has not been detected in schizophrenia brain tissue, and indeed, many studies searching for cerebral viral or bacterial DNA or protein postmortem have been negative. However, as already suggested, given the precocious timing of certain infections related to schizophrenia risk, an inability to detect viral traces many decades later at autopsy is perhaps not unexpected. A viral presence in cerebral tissue might also not be strictly necessary, and several groups have suggested that circulating infection-related cytokines may participate in the risk-promoting process. For example, maternally administered interleukin (IL)-6 or the cytokine releaser polyriboinosinic-polyribocytidylic acid are also known to be able to influence the behavior of the adult offspring in rats and also induce morphological changes in the hippocampus resembling those observed in schizophrenia. Immune perturbation can also affect behavior in adults. For example, in adult mice, T-cell deficiency can induce cognitive deficits, while T-cell vaccination can reduce the behaviorally excitant and cognitive-disturbing effects of the NMDA receptor antagonist MK-801 or amphetamine. Both heat shock (hyperthermia/fever) and cytokines (interferons, IL1B, and TNF) are able to activate the viral-activated eif2alpha kinase pkr. (EIF2AK2: eukaryotic translation initiation factor 2-alpha kinase 2) involved in the control of protein translation (see above). Infection or its inflammatory or immune consequences may, thus, affect both neurodevelopment and neurotransmitter-related behavior, although the mechanisms behind these effects remain poorly understood. Again, as with individual susceptibility genes, the implication of individual pathogens as risk factors has been subject to controversy.

Pathogens purloin different aspects of the hosts’ physiology during their life cycles. They have evolved mechanisms to adhere and gain entry to target cells via specific receptors and use endocytic pathways and the intracellular transport and sorting machinery to access specific cellular compartments. They may use cellular junctions for intercellular spread, or lie dormant in vacuoles or vesicles, formed by host and pathogen proteins. Viruses usurp the host’s transcriptional machinery to replicate. Each pathogen uses its own and the hosts’ cellular genes and proteins during these processes. The host retaliates using the immune system, as well other infection-related defense mechanisms, including cytokine production and free radical–related bactericidal and viricidal effects. Pathogens have also evolved mechanisms to avoid immune detection or to otherwise abrogate immune defense.

As described below, many genes associated with schizophrenia in genetic studies code for proteins specifically implicated in this host-pathogen battlefield creating a further network of genes, in this case etching out stages of the pathways used by the pathogens during their life cycles and interaction with the host. These extrinsic pathways may play a role in infection, although the host genes involved evidently also play a role in human physiology that is often pertinent to schizophrenia. Again, multifunctional effects seem to dictate the relevance of the gene as certain genes of high interest, notably, DISC1, NRG1, RGS4, and dysbindin, inter alia, can be related to both intrinsic and extrinsic signaling networks. This might suggest that association studies may in some cases relate to genes specifically involved in the host/pathogen physiology of causative pathogens as well as to the disease itself. If these genes are implicated in the life cycles of the pathogens, polymorphisms therein may well affect the virulence of the pathogen, providing a further example of how genes and environmental factors might interact to produce
disease. These interactions have important medical implications and suggest that targeting the appropriate pathogens may have dramatic effects on the incidence and progression of schizophrenia as already suggested by several authors (for review, see Yolken and Torrey\(^7\)).

Methods and Background

The genes associated with schizophrenia have been the subject of previous reviews\(^2,22\) and are listed together with the environmental risk factors at http://www.polygenicpathways.co.uk. The pathogens included in this review focus on influenza, rubella, herpes viruses (HSV-1 and CMV), or the protozoan parasite \textit{T gondii}. A literature survey was undertaken to analyze whether any of the susceptibility gene candidates are associated with the cellular processes used during the life cycles of these and other potential pathogens. The Vita database (http://vita.mbc.nctu.edu.tw/)\(^{28}\) was used to predict potential viral targets of human microRNAs. Because of the large volume of data, the bulk of this analysis, and references, are posted in a supplementary table at http://www.polygenicpathways.co.uk/schizmicrobes.htm, and only certain main conclusions are summarized below. The families of genes mentioned in this review are annotated in a supplementary table posted at http://www.polygenicpathways.co.uk/szfam.htm. Association data relating to genes implicated in infection in man are summarized at http://www.polygenicpathways.co.uk/infectassoc.htm. Genes associated with schizophrenia are in \textbf{bold} throughout the text.

Viral Receptors

\textbf{Binding to Chemical Residues on the Cell Surface (figure 1).} Influenza, herpes simplex, and CMV bind to sialic acid residues on the host cell surface, including alpha-2-8-linked sialic acids synthesized by the sialyltransferase \textit{SIAT8B}.\(^{29}\) Acetylated galactosamine (metabolized by \textit{GALNT7}) is also a receptor for the influenza C virus glycoprotein,\(^{30}\) while glycoprotein C of the herpes virus acquires N-acetylgalactosamine prior to routing to the host Golgi apparatus.\(^{31}\) The CMV binds to sulfated glucuronyl glycosphingolipids (synthesized by \textit{B3GAT1}).\(^{32}\) The rubella virus binds to membrane phospholipids and glycolipids, and phospholipase A2 (\textit{PLA2G4}) digestion of Vero cells markedly reduces viral infectivity. Viral infection is also reduced after beta-N-acetyl-d-glucosaminidase, alpha-glucoosidase, and beta-galatosidase treatment, suggesting that diverse carbohydrate residues may contribute to a complex receptor structure for this virus.\(^{33}\) Aspartylglucosaminidase (\textit{AGA}) metabolizes beta-N-acetyl-d-glucosamine residues.

\textbf{Ligand-Activated Receptors (figure 1).} The herpes virus uses a number of receptors to gain entry to host cells,\(^{34}\) including nectin receptors (PVRL1-3), for which \textbf{PICK1} acts as a scaffold\(^{35}\); a TNF-related receptor (\textit{TNFRSF14}), for which lymphotoxin-alpha (\textit{LTA}) is the natural ligand; and the fibroblast growth factor receptor 1 (\textit{FGFR1}).\(^{36}\)

The epidermal growth factor (\textit{EGF}) receptor is used by CMV for entry, and epidermal growth factor receptor (\textit{EGFR})/ERBB3 heterodimers bind glycoprotein B of the virus.\(^{37}\) \textit{NRG1} is a ligand for such heterodimers.\(^{38}\) While \textit{RGS4} is involved in ERBB3 signaling,\(^{39}\) A CMV viral homolog of the cytokine \textit{IL10} also binds to the host cognate receptor \textbf{IL10RA}.\(^{40}\)

\textbf{Adhesion Molecules and Cell Junctions.} Although none of the adhesion molecules (\textit{CHL1}, \textit{NCAM1}) or junctions (\textit{CLDN5}, \textit{GJA8}) implicated in schizophrenia are specifically involved in the life cycles of the pathogens in this study, these families play an important role in microbe recognition (adhesion molecules)\(^{41}\) and intercellular spread (gap and tight junctions).\(^{42}\) For example, \textit{NCAM1} serves as a viral receptor for the herpes family rabies virus,\(^{43}\) while claudins function as entry receptors for hepatitis C.\(^{44}\) \textit{ARVCF} is a component of adherens junctions, which play an important role in the invasion of several viruses including herpes simplex.\(^{45}\) The chitinase-like protein, \textit{CHI3L1}, is also involved in bacterial adhesion and invasion in colonic epithelial cells.\(^{46}\)

\textbf{Intracellular Traffic.} Many pathogens exploit the trafficking networks used by the cell to obtain nutrients, or to endocytose membrane receptors, entering cells via these routes (figure 1). These endosomal and other pathways converge on lysozymes, which are able to kill pathogens.\(^{47}\) Dysbindin (\textit{DTNBP1}), \textit{BLOCIS3}, or \textit{MUTED} are part of the biogenesis of lysosomal organelles complex (BLOC-1) which is specifically involved in the sorting of cargoes from vacuolar early endosomes to the lysosomal complex.\(^{48}\) Viruses also use the microtubule network to gain access to the nucleus and are driven along by the host’s dynein and kinesin motors toward the microtubule organizing center close to the nucleus.\(^{49}\) Viral inclusions, factories for viral replication, form at pericentriolar sites close to this microtubule organizing center or in specialized nuclear domains known as ND10/PML bodies.\(^{50}\) \textit{DISC1} is a component of this microtubule-related dynein motor complex\(^{51}\) and implicated in pericentriolar function via its association with kendrin, a binding partner of the pericentriolar protein \textit{PCMi}.\(^{52}\) While neither \textit{DISC1} or \textit{dysbindin} nor the kinesin or microtubule genes (\textit{KIF2A}, \textit{MAP6}, \textit{TUBA2}) have been specifically implicated in viral traffic, they are clearly involved in processes commandeered for viral transport.

The phosphoinositide kinase subtype \textit{vps34} (\textit{PIK3C3}) is involved in the vacuolar protein-sorting pathway used by the influenza virus to gain entry to cells. Phosphoinositide-5 and -4 kinases (\textit{PIPK2A}, \textit{PI4K}) are also involved in the intracellular traffic of the influenza virus.\(^{53}\)
Protein Folding and Heat Shock Proteins. Proteins derived from the influenza, CMV, herpes simplex, and rubella all bind to calreticulin (CALR), which plays an important role in viral protein folding. The CMV glycoprotein B binds to a number of heat shock proteins including HSPA1L and HSPA1B, also a binding partner of the influenza ribonucleoprotein and polymerase complex. Subjugation of the host’s protein folding machinery by viral polypeptides also leads to endoplasmic reticulum stress. ATF6 and XBP1 play a key role in this pathway (see below and figure 4).

Viral Replication

Nuclear Import. Many viruses need to enter the cell nucleus, where they subvert the cellular transcriptional machinery (figures 1 and 2). Both RANBP5 and KPNA3 (nuclear importin or karyopherins that control the entry of proteins from the cytoplasm to the nucleus via the nuclear pore) are involved in the nuclear import of the influenza virus.

RNA Polymerase and Viral Transcription. Herpes simplex uses the host’s RNA polymerase II, which it diverts to synthesizing viral RNA at the expense of the host’s (figure 1). The mediator complex (MED12 [aka, HOPA], MED15 [aka, PCQAP]) is a host RNA polymerase coactivator used by herpes simplex, which diverts the polymerase activity toward viral RNA synthesis. A viral protein VP16 binds to components of this mediator complex (MED15, MED17, and MED25). Other schizophrenia susceptibility candidates also affect the transcription of the influenza virus. Vitamin B6, the product of pyridoxamine 5’-phosphate oxidase (PNPO), inhibits influenza viral replication by a direct action on the viral transcriptase to which it binds. NRG1 signaling via ERBB receptors might also be expected to influence influenza viral transcription. A host protein, EBP1, binds to and acts as an inhibitor of the influenza virus RNA polymerase. EBP1 was initially characterized as an ERBB3 binding partner. It dissociates from this receptor following stimulation by its ligand, NRG1, and translocates from the cytoplasm to
the nucleus. RGS4 is a component of the ERBB3 signaling pathway. The effects of NRG1 on influenza infection have not been directly examined, although interestingly, NRG1 expression levels are markedly increased in the blood of influenza-infected children with febrile convulsions, while RGS4 expression levels are increased in the brains of influenza-infected mice.

Viral RNA Methylation: S-adenosylmethionine and Homocysteine Metabolism. Viral RNA methylation, using the methyl donor S-adenosylmethionine, is crucial for the maturation of viral RNA as shown in figure 2 (modified from a review by De clerq) (figure 2). The product of these methylation reactions, S-adenosylhomocysteine inhibits the diverse methyltransferases involved in this process. S-adenosylhomocysteine (SAH) is normally rapidly metabolized to adenosine and homocysteine by SAH hydrolase (ADCY), and inhibitors of this enzyme inhibit viral RNA methylation by favoring SAH accumulation. SAH hydrolase inhibitors (carbocyclic adenosine, carbocyclic 3-deazaadenosine, neplanocin A, 3-deazaneplanocin A) display broad-spectrum antiviral activity in the laboratory that has not yet translated to the clinic, presumably, because of the difficulty of selectively affecting viral rather than host methylation.

Methyltransferase inhibitors also inhibit influenza viral replication, and methylation inhibition also results in the increased nuclear retention of influenza viral RNA. Methylation of viral DNA or proteins also plays a role in the viral life cycles. For example,

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**Fig. 2.** Viral RNA Methylation. Using S-adenosylmethionine, Plays an Important Role in Viral RNA Maturation. The genes involved in S-adenosylmethionine and homocysteine metabolism are depicted here. Gray-shaded boxes represent those associated with schizophrenia (modified from De clerq).
HIV-1 latency or lytic infection can be influenced by such reactions.66 The availability of S-adenosylmethionine is, thus, a general factor contributing to viral viability. One might also expect that methylation of viral RNA would be a competing factor for the methylation of host DNA or protein, depending on the compartment in which this occurs. Viruses also affect the methylation of host DNA. For example, the herpes simplex strain HSV-2 produces marked hypomethylation of host DNA in infected rat embryo cells, although this effect may be due to a redistribution of the host DNA methyltransferase rather than to any diversion of methionine/homocysteine metabolism toward viral methylation.67 A generalized decrease in DNA methylation has been observed in leukocytes isolated from male schizophrenic patients.68 Several genes implicated in schizophrenia (MTHFR, MTR, and MTHFD1) are relevant to this area as shown in figure 2. Other methylating enzymes may also influence this cycle indirectly by modifying substrate levels. For example, phosphatidylethanolamine methyltransferase (PEMT) knockout mice have dramatically reduced plasma homocysteine levels69 and catechol-O-methyltransferase (COMT) inhibition increases S-adenosylmethionine levels in the rat brain.70 DNA methyltransferase expression is markedly increased in certain areas of the schizophrenic brain, and S-adenosylmethionine levels are also increased. Such effects have been proposed to be relevant to the hypermethylation of the reelin promoter and a reduction in its messenger RNA (mRNA) and protein levels observed in schizophrenia.71–73 While not wishing to imply that viruses are fully responsible for these effects, or that the effects of genes such as COMT are solely related to viral replication, it is clear that viruses are able to affect host methylation reactions and that these in turn are able to influence viral replication. Interestingly, the prenatal infection of mice with influenza does result in a reduction in cortical and hippocampal reelin immunoreactivity.74

**Human microRNA**

MicroRNAs are short RNA transcripts, derived from chromosomal DNA, that are not translated into protein (figure 1). Their sequence is complementary to one or more mRNAs, to which they bind, thus, inhibiting translation of the mRNA to protein. They can bind either to host mRNA or to viral genomes, wherever sequence complementarity exists.

The human microRNAs, hsa-mir-206 and hsa-mir-198, are complementary to a number of host mRNAs and also to several viral RNAs, as suggested by interrogation of the Vita database.4 This bioinformatic analysis suggested that hsa-mir-198 targets viral genes encoding for the influenza virus PB2 protein and for structural proteins E1, E2, and C of the rubella virus, inter alia. hsa-mir-206 is predicted to bind to genes of the human coxsackievirus B2 and human poliovirus 1, inter alia.

**Bacteria and Parasites**

This review is primarily concerned with viral risk factors, although it is likely that such interactions are also pertinent to the life cycles of bacteria (eg, Chlamydia) or parasites (T gondii) that have been implicated in schizophrenia (figure 3). Only a few examples are given here. Tryptophan, metabolized by tryptophan hydroxylase (TPHI)-1, is an essential nutrient for several microorganisms, including T gondii and chlamydial species.75

Spirochetes (T gondii) forage phospholipids in the blood. They possess an enzyme that metabolizes dihydroxyacetone phosphate (glycerone-3-phosphate) to glycerol-3-phosphate that may be used as an energy source by the parasite. Glyceronephosphate O-acetyltransferase (GNPAT) metabolizes glycerone-3-phosphate to acylglyceronephosphate.76

Exogenous cyclic adenosine monophosphate (AMP) also stimulates T gondii growth in cell culture and cyclic AMP phosphodiesterase inhibition or activation and stimulates or inhibits T gondii growth, respectively77 (cf, the cyclic nucleotide phosphodiesterases PDE4B and CNP1). Toxoplasma gondii, like viruses, also binds to siaic acid residues on cell surfaces78 (cf, SIA18B). Toxoplasma gondii contact with the cell surface in fibroblasts is mediated by laminins on the parasite surface, which bind to host beta 1 integrins.79 Interestingly, the reelin/beta 1 integrin system controls the surface expression of NMDA receptors (GRIN1, GRIN2B),80 although the effects of T gondii on reelin-NMDA receptor interactions have not been studied. A T gondii cyclophilin-like secreted protein also binds to the chemokine receptor CCR5.81 The parasite survives in macrophages by hiding in parasitophorous vacuoles that avoid fusion with lysosomes. This parasitophorous vacuole is derived from invagination of the host plasma membrane following entry of the parasite by active penetration. The parasite uses the host’s microtubule network to transport host endosomes or lysosomes to invaginations within this organelle and feeds on their constituents. This vacuole is also closely associated with the host mitochondria and endoplasmic reticulum,82 and its parasite-derived constituents dense granule proteins have been shown to bind to a number of host proteins including the transcriptional regulator PAX6 and the quinone reductase NQO2.83 Activation of the immune response can reroute these vacuoles to the lysosomal system allowing macrophage bactericidal activity. The phosphoinositide kinase PIK3C3 plays a key role in this vacuolar/lysosomal fusion and in enabling macrophages to kill T gondii.84 The cellular microtubule network and cell-sorting machinery, thus, play an important role in both viral and parasitic infection.
General Pathways

Growth Factor and Phosphoinositide Signaling. Many pathogens subvert the phosphoinositide kinase/AKT survival pathway, primarily to activate antiapoptotic pathways that allow the pathogen to stay in residence (figure 4). The NS1 influenza viral protein binds to the p85 beta subunit of phosphoinositide kinase (PIK3R2) and activates AKT1 signaling. These effects limit the viral-induced cell death program allowing further replication. CMV, herpes simplex, rubella, and T gondii also use this type of subversion of growth factor signaling (see supplementary data on Web site).

Glutathione and Oxidative Stress. Glutathione levels are decreased in the cortex and cerebrospinal fluid (CSF) in schizophrenia (figure 4). Glutathione levels are also...
decreased in Vero cells by HSV-1 infection, and glutathione supplementation markedly inhibits viral replication.\textsuperscript{88} HSV-1 infection of microglia results in the upregulation of the glutamate/cysteine transporter \textit{SLC1A2}, which plays an important role in providing the glutamate and cysteine necessary for glutathione biosynthesis by the glutamate cysteine ligases (\textit{GCLC}, \textit{GCLM}). High intracellular glutathione levels also inhibit the replication of the influenza virus,\textsuperscript{89} and cells with high levels of glutathione are also resistant to CMV infection.\textsuperscript{90} Glutathione is

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\textbf{Fig. 4.} Oxidative Stress, Endoplasmic Reticulum Stress, Starvation, Viruses, Bacteria, and Other Pathogens Activate the eif2alpha kinase Signaling Network. Phosphorylated eif2alpha shuts down protein synthesis via inhibition of the translation initiation factor eif2B, a gene that is crucial to oligodendrocyte function. Protein synthesis is also controlled by growth factors via the \textit{AKT1} signaling network and by NMDA and metabotropic glutamate receptor activation. Protein synthesis is enhanced by long-term potentiation and repressed by long-term depression via effects on eif2alpha phosphorylation. \textit{ATF4} activation by this pathway regulates processes designed to counteract the effects of these stressors, or in unsuccessful, to trigger apoptotic programmes. Pathogens activate the eif2alpha kinase pkr or Toll receptor signaling and can activate or inhibit the AKT1 survival pathway. Gray boxes depict schizophrenia susceptibility genes identified in association studies or as genes within runs of homozygosity regions marked with an asterisk (\textit{ATF6}, eif2alpha/\textit{EIS2S1}). Those implicated specifically in the life cycles of each pathogen are also illustrated. Hatched boxes represent viral proteins or genes, which interact directly with certain of the proteins in this network (see text for details). LTP, long-term potentiation; LTD, long-term depression; HSV, herpes simplex virus; CMV, cytomegalovirus.

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also effective in reducing influenza infection in mice in vivo.91

NMDA receptors possess a redox modulatory site sensitive to glutathione, and glutathione depletion in animal models leads to NMDA receptor hypofunction, mimicking several features of schizophrenia (reviewed in Lavoie et al92). A recent clinical trial using the glutathione precursor N-acetylcysteine has reported significant modifications in auditory evoked potential deficits in schizophrenia and also suggested effects on the improvement of negative symptoms.92 As glutathione-based therapy might be expected to reduce viral load and favorably modify NMDA receptor function, this area clearly deserves further attention. Glutathione also induces the egress of T gondii from infected cells by activating a T gondii–secreted apyrase (nucleoside triphosphate hydrolase) in the parasite vacuole resulting in a rapid depletion of host cell adenosine triphosphate (ATP).90

The Innate Immune System. A large number of cytokine- and chemokine-related genes (CCR5, CSF2RA, CSF2RB, IL1B, IL1RN, IL2, IL3, IL3RA, IL4, IL10, IL10RA, IL12B, IL18, IL18R1, IL18RAP, LTA, TNF) and major histocompatibility complex antigens (HLA-A10, HLA-B, HLA-DRB1) have been implicated in schizophrenia (figure 4). The angiotensin-converting enzyme is also involved in immune processes and cleaves an influenza antigen nucleoprotein.93 Tripeptidyl peptidase 2 is also involved in the antigen processing of the influenza virus.94 A CMV protein (UL16) also binds to the natural killer cell receptor ligand MICB. This receptor triggers natural killer cells and costimulates antigen-specific T cells that would normally destroy invading pathogens. MICB binding by UL16 may circumvent this immune response.95 The CMV IE2 protein also binds to the promoter of the IL2 gene and upregulates its expression.96 Free radical–generating systems also play an important role in pathogen defense. For example, nitric oxide synthase 1 (NOS1) is toxoplasmacidal.97

The immune system is designed to protect against pathogens by recruiting immunocompetent cells, macrophages, and microglia to sites of infection, and in a general sense, polymorphisms in many of these genes would by expected to impinge upon the life cycles of any of the pathogens implicated in schizophrenia. Most of these genes are affected in some way by one or other of the pathogens and several are able to influence pathogen viability. The specific relationships of these genes to each of the pathogens are reviewed and referenced more extensively in the supplementary table at http://www.polygenicpathways.co.uk/schizmicrobes.htm, and their overall effects summarized in table 1. (A number of other genes whose function one would tend to associate with central nervous system (CNS) effects are also expressed on lymphocytes and play a role in the immune system. These include the growth factors BDNF and NTF3, NMDA and metabotropic glutamate receptors, and dopamine and serotonin receptors, inter alia. The effects of polymorphisms on immune function may be as important as their effects in the CNS. Further details are provided at http://www.polygenicpathways.co.uk/schizmicrobes.htm.)

Glutamate Receptors and Plasticity. A large proportion of the genes implicated in schizophrenia code for presynaptic and postsynaptic proteins related to glutamatergic neurotransmission and to components of the postsynaptic density (figures 1 and 3). Few of these appear to be directly implicated in the life cycles of the pathogens implicated in schizophrenia, although both FYN and citron kinase, binding partners of the herpes simplex and rubella virus, respectively, are components of the postsynaptic density, while PICK1 is a scaffolding partner for glutamate receptors and the herpes entry portals, nectin receptors35 (figure 1). The binding of T gondii species to integrins, components of the reelin signaling pathway, might also be expected to influence NMDA receptor function (figure 2). The NMDA receptor does bind to components of the HIV virus,98 and the use of this receptor by other viruses or pathogens would not be unprecedented.

Viral infection does affect glutamatergic signaling. For example, the influenza nucleoprotein distributes to dendritic spines in cultures of rat hippocampal neurones. Spontaneous excitatory synaptical activity and the amplitude of miniature excitatory postsynaptic currents are reduced at later stages, 17–22 days after transfection.99 Infection of PC12 cells with the herpes virus HSV-1 also results in a reduced expression of the NMDA receptor subunit GRIN1.100 The hippocampal expression of GRIN1 is also reduced following CMV infection in mice, an effect also observed in primary neuronal culture.101

Interestingly, the NS1 influenza protein binds to the RNA transport protein staufen102 which plays a key role delivering mRNA to neuronal dendrites. Dendrites contain mRNA, thought to code for many of the elements necessary for glutamatergic transmission, protein synthesis, and plasticity.103 Staufen plays a key role in dendritic mRNA transport, allowing delivery of these genes.104

The herpes virus also triggers the formation of varicosities along the axons of sensory neurones that stain for synaptophysin and closely resemble presynaptic terminals.105 These effects are mediated by viral activation of the rho guanosine triphosphate–binding protein CDC42, which also plays a key role in the maturation of presynaptic glutamatergic terminals.106 These varicosities serve as exit sites for the herpes virus, and if they correspond to presynaptic terminals, presumably, contain many of the components of these structures (eg, synapsins, complexins, and syntaxins). CDC42 also controls...
Table 1. A survey of the Interactions Between the Pathogens and Genes Implicated in Schizophrenia

<table>
<thead>
<tr>
<th>Pathogen</th>
<th>Influenza</th>
<th>Cytomegalovirus</th>
<th>Herpes</th>
<th>Rubella</th>
<th>Toxoplasma</th>
<th>Others</th>
</tr>
</thead>
<tbody>
<tr>
<td>Direct interactions (binding, degradation, or synthesis of a binding partner)</td>
<td>ACE</td>
<td>CALR</td>
<td>APOE</td>
<td>CALR</td>
<td>CCR5</td>
<td>ARVC/Herpes</td>
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<td>Herpes</td>
<td>Rubella</td>
<td>Toxoplasma gondii</td>
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<td><strong>Kills pathogen or reduces replication/infection</strong></td>
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<td>Glutathione (GCLM, GCLC, GSTM1, GSTT1)</td>
<td>Glutathione (GCLM, GCLC, GSTM1, GSTT1)</td>
<td>Glutathione (GCLM, GCLC, GSTM1, GSTT1)</td>
<td>CCR5del</td>
<td>NRG1 releases EB1, an influenza viral transcriptase inhibitor,</td>
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<td><strong>% of schizophrenia candidates to date (N = 245)</strong></td>
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<td><strong>Genes implicated in infection</strong></td>
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<td>(association studies related specifically to these pathogens)</td>
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<td>HLA-DRB1, IL10</td>
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<td>No reports of association with schizophrenia</td>
<td>No reports of association with schizophrenia</td>
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<td>% of genes implicated in infection also implicated in schizophrenia Immune defense</td>
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<td>2 of 3 = 66% (CCR5, CSF2RA, CSF2RB, IL1B, IL1RN, IL2, IL3, IL3RA, IL4, IL10, IL10RA, IL12B, IL18, IL18R1, IL18RAP, LTA, TNF), HLA antigens (HLA-A10, HLA-B, HLA-DRB1), and genes involved in antigen processing (TPP2) or T- or B-cell signaling and development (BDNF, CTLA4, EGR2/3/4, FOXP2, GRIN1, GRIN2B)</td>
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**Note:** Most of the interactions summarized here are referenced in a supplementary table posted at http://www.polygenicpathways.co.uk/schizmicrobes.htm. Association data with respect to infection per se are posted at http://www.polygenicpathways.co.uk/infectassoc.htm. See text for details.
dendritic spine development,\textsuperscript{107} a process disrupted in the schizophrenic brain. The M1 protein of the influenza virus also binds to CDC42,\textsuperscript{108} whose suppression inhibits viral production.

**Dopaminergic and Serotonergic Systems.** Dopamine or serotonin receptors have not been implicated in the life cycles of any of the pathogens detailed in this review, although the 5HT-2 receptor (HTR2A) is an entry portal for the JC polyoma virus that causes demyelination (figure 3).\textsuperscript{109} Pathogenic infection does, however, have marked effects on serotonin metabolism. The activation of pathogen recognition receptors (the viral-activated kinase pkr and Toll receptors) stimulates the production of interferons and other type 1 cytokines (IL1B, TNF) involved in immune defense. These stimulate the expression and activity of the tryptophan-degrading enzyme, indoleamine 2,3-dioxygenase (INDO). Type 2 cytokines (eg, IL4, IL10) inhibit INDO activity. Such effects modify the concentrations of the serotonin precursor, tryptophan, and the enzyme product, N-formylkynurenine in different cellular compartments. Tryptophan is an essential nutrient for several microorganisms, including \textit{T gondii} and chlamydial species,\textsuperscript{75} while N-formylkynurenine is a precursor of kynurenic acid, an endogenous antagonist of both nicotinic, and NMDA receptors. Kynurenic acid levels are elevated in the CSF of schizophrenic patients, and as already reviewed, such effects may well be due to the activation of inflammatory cascades by invading pathogens and could well contribute to the neurotransmitter imbalances at the root of schizophrenic symptoms.\textsuperscript{110} Tryptophan depletion also exerts antiviral effects against herpes strains, suggesting a role of the INDO pathway in general immune defense.\textsuperscript{75} It, thus, appears that infection, or its consequences, can have profound effects on the glutamate/dopamine-serotonin triad that plays such a significant role in the symptomatology of schizophrenia.

**Antigenic Molecular Mimicry.** Molecular mimicry has been reported between the myelin-oligodendrocyte protein (MOG) and structural glycoprotein E(2)of the rubella, or CMV-UL86 peptide, between a CMV envelope glycoprotein H peptide and tyrosinase (TYR), and between glutamat decarboxylase 1 (GAD1) and a rubella antigen in diabetic patients (figures 1 and 3). CMV infection correlates with immunoglobulin M anti–myelin-associated glycoprotein (anti-MAG)/sulfated glucuronyl paragloboside antibody levels in neuropathic conditions. Mimicry has also been reported between the NMDA receptor subunit GRIN2B and the influenza virus hemagglutinin (see Web site for references).

**The eif2alpha Kinase Signaling Network.** Viruses activate the eif2alpha kinase pkr (EIF2AK2), which phosphorylates eif2alpha (figure 4). This leads to an inhibition of protein translation because phosphorylated eif2alpha inhibits the actions of the translation initiation factor eif2b. Other stressors including starvation and oxidative and endoplasmic reticulum stress also activate this pathway via the stimulation of other eif2alpha kinases specific for each stressor. The protein translation pathway is also regulated by growth factors (eg, BDNF, EGF, FGF1, NRG1) and by glutamate receptors (eg, GRIN1, GRIN2A). Oligodendrocytes are particularly sensitive to eif2b malfunction because mutations in the genes coding for subunits of this complex cause vanishing white matter disease.\textsuperscript{3} The malfunction of this signaling network has been suggested as a potential cause of the oligodendrocyte cell death observed in schizophrenia.\textsuperscript{111} eif2Alpha phosphorylation also activates the transcription factor ATF4 (a binding partner of DISC1), whose outputs control a series of processes designed either to rectify the initial effects of the stressors (eg, the activation of oxidative stress–related or protein folding–related pathways in response to oxidative or endoplasmic reticulum stress).\textsuperscript{3}

The viral pathogens implicated in schizophrenia activate the viral-sensitive kinase pkr and \textit{T gondii} as well as bacterial pathogens can be linked to this network via their effects on Toll receptors. Toll-like receptors play an important role in activating the immune response defense mechanisms in response to viruses, bacteria, and other pathogens. Their signaling networks also activate the double-stranded DNA-responsive eif2alpha kinase pkr (EIF2AK2).\textsuperscript{112}

A number of herpes viral products affect components of this pathway. For example, the HSV-1 gene ICP34.5 encodes for a phosphatase that mimics the actions of gadd34 (PPP1R15A) whose normal role is to dephosphorylate eif2alpha. The herpes protein US11 prevents pkr (EIF2AK2) activation by inhibiting its activator protein kinase, interferon-inducible double stranded RNA dependent activator (PRKRA).\textsuperscript{113} The herpes viral glycoprotein B (UL27) also binds to perk (EIF2AK3), preventing its activation by endoplasmic reticulum stress.\textsuperscript{114} The CMV also activates perk (EIF2AK3), independently of its effects on the viral kinase pkr, and induces splicing of XBP1 to its active form.\textsuperscript{115} The CMV protein IE2 also binds to nrf1 and nrf2 (NFE2L1 and 2),\textsuperscript{116} the transcription factors that control mitochondrial biogenesis and glutathione-related genes.

The latency transcript of the HSV is a promoter region of the viral genome responsible for controlling the reactivation of the virus. It contains 2 cyclic AMP response elements, which bind to cAMP response element-binding (CREB) and ATF4 (also known as CREB-2),\textsuperscript{117} suggesting that disruption of this signaling network may also affect viral latency. The human CMV major immediate-early enhancer also contains 5 functional cyclic AMP response elements and activation of the cyclic AMP signaling cascade by forskolin is able to modify the function of

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\textsuperscript{3} This manuscript was downloaded from http://schizophreniabulletin.oxfordjournals.org/ on May 1, 2016.
certain of the features of schizophrenia in mice. It is of course possible to mimic certain of the symptoms of schizophrenia without advocating a role for exogenous pathogens. For example, amphetamine or phencyclidine can reproduce certain of the features of psychosis.\textsuperscript{120} DISC1 mutant mice also display a range of schizophrenia-like phenotypes, including ventricular enlargement, reduced parvalbumin immunoreactivity in cortical interneurons, hyperactivity, and modified sensorimotor gating.\textsuperscript{121} Disruption of ERBB4 signaling can also lead to hypomyelination and to subsequent dopaminergic hyperactivity in mice.\textsuperscript{122} These features (endophenotypes) could, thus, be induced in man by polymorphisms in any or several of the multiple susceptibility genes implicated in NMDA receptor function and plasticity, or in those involved in NRG1 or DISC1 signaling. The pathways carved out by such genes may well form an intrinsic network whose dysfunction may contribute to much of the schizophrenia pathology.

Prenatal infection or immune activation can also model certain of the features of schizophrenia in mice.\textsuperscript{123} The pathogens implicated in schizophrenia affect the same signaling network, the eif2alpha kinase pathway, as that influenced by many of the susceptibility genes, and as described above, interact in multiple ways with a number of gene candidates. Because these pathogenic and genetic pathways coalesce, it is possible to advocate some form of synergy between pathogenic and susceptibility genes and pathways.

Others genes, eg, the immune-related genes, those coding for receptors (EGFR, FGFR1, IL10RA), or metabolizing compounds (AGA, B3GAT1, SIA1TB8, GALNT7, PLA2, PSPA) that these pathogens bind or forage (GNPAT, TPH1), those involved in the nuclear/cytoplasmic transport of the influenza virus (KPN3, RANBP5), or implicated in RNA polymerase function (MED12, MED15, ZNF74) may trace out pathways more related to the life cycles of the pathogens implicated in the disease than to the disease itself. In many cases, dual functions may explain the greater importance of a particular gene. For example, NRG1 signaling affects glutamatergic signaling and oligodendrocyte function and might also be expected to affect influenza virulence via its effects on the viral transcriptase inhibitor EBP1.\textsuperscript{50,61} COMT might be expected to modify dopaminergic function but could also affect viral methylation via its effects on S-adenosylmethionine levels. The involvement of DISC1 in the control of the microtubule network suggests that it might be important both in viral traffic and in the rerouting of microtubules to the vacuoles formed by T. gondii, as well as in important aspects of neurobiology. DISC1 also binds to ATF4 that plays such a key role in the stress- and pathogen-activated eif2alpha defence pathway. Currently, these ideas are speculative as the effects of NRG1; COMT or DISC1 variants on viral or parasitic virulence have not been studied. It is known, however, that several immune defense genes implicated in schizophrenia are also associated with infectious agents (see below).

Integrated Pathways: Genetic and Pathogenic Units Contributing to Disruption of the Same Networks. Schizophrenia is a complex disorder with multiple subpathologies or endophenotypes, which broadly match the gene families implicated in the disorder. For example, subsets of genes can be related to glutamatergic function, oligodendrocyte viability, neuronal growth and development, oxidative stress or dopaminergic, serotonergic, or GABAergic function, inter alia. These genes form a number of integrated networks (see “Introduction”). While certain important “hub genes,” eg, DISC1, appear to bridge these distinct families and networks, it seems currently unlikely that any single gene defect is able to provoke all the underlying pathologies that constitute the entire complexity of schizophrenia. The full-blown pathology is, thus, likely to relate to an integration of multiple phenotypes each related to their corresponding underlying genotypic subgroups.

When pathogen genes or proteins interact with components of these signaling networks, regardless of whether they interact with the susceptibility genes themselves, they essentially become a part thereof and could be regarded simply as yet another susceptibility unit (albeit foreign) capable of disrupting the same pathways carved out by the human susceptibility genes (see figures 1–4). The ability of the influenza virus to interfere with staufen or cdc42 could clearly affect both presynaptic and postsynaptic glutamatergic networks, while the effects of many pathogens on the PI3K/AKT survival pathway could evidently interfere with multiple growth factor signaling networks involved in plasticity or survival. Certain viruses are known to modify synaptic activity. Both herpes simplex and murine CMV infection results in the downregulation of NMDA receptor expression in PC12\textsuperscript{100} cells or in the hippocampus.\textsuperscript{101} In neuronal/glial coculture, the infection of murine astrocytes by CMV increases their sensitivity to ATP and glutamate, while the noninfected overlying neurons exhibit decreased synaptic activity as a result.\textsuperscript{124}
Because each pathogen is able to activate the pkr signaling network that controls translation initiation (eif2B), that in turn appears to be crucial to oligodendrocyte viability and to synaptic plasticity, it seems plausible that compromised oligodendrocyte and synaptic function, as observed in schizophrenia, might be related to such pathogens. Indeed, prenatal influenza infection in mice results in white matter thinning in the corpus callosum\textsuperscript{125} as well as atrophy of cortical pyramidal cells.\textsuperscript{126} Oligodendrocyte cell loss or demyelination has also been observed in response to infection with rubella\textsuperscript{127} or herpes simplex in vitro.\textsuperscript{128} The CMV appears to preferentially infect human astrocytes and neurons, although oligodendrocyte infection has been reported. Interestingly, in a human oligodendrocyte cell line, resistance to CMV infection can be overcome by treatment of the cells with the protein kinase C activator, phorbol 12-myristate 13-acetate, or with a combination of dibutyryl cyclic AMP and a phosphodiesterase inhibitor (cf, \textit{PDE4B}, \textit{CNP}), suggesting that cellular activity can modulate the infection of such cells.\textsuperscript{129} Other schizophrenia-related endophenotypes induced by these pathogens include defects in prepulse inhibition in the offspring of mice infected with the influenza virus,\textsuperscript{130} while in adult mice, influenza infection results in decreased anxiety and impaired spatial learning.\textsuperscript{16} Thus, viruses, per se, can impact on several endophenotypes involved in schizophrenia. Prenatal immune challenge in mice also leads to dopaminergic-related hyperactivity and cognitive impairments in the adult progeny,\textsuperscript{123} although the mechanisms leading to such effects are poorly understood.

As with each susceptibility gene, it seems unlikely that any single pathogen could produce all the pathological features of schizophrenia per se, but each may affect certain signaling networks, related to different susceptibility gene subsets, that in turn impact on particular endophenotypes.

\textit{Comparative Figures in Relation to Other Gene Families and Pathological Processes.} Viruses and pathogens interact with many human genes and proteins during their life cycle, and by chance, one would expect any large gene data set to contain a number of pathogen-interacting elements. To be able to define any overall statistical relevance, one would need a comprehensive data set of all the pathogen-host interactions for each pathogen. While several protein interaction databases exist for human proteins, such data sets are not currently available for human-pathogen interactions for the pathogens reviewed in this study. Perhaps, the most intensively studied virus is the Acquired immune deficiency syndrome (AIDS) human immunodeficiency virus (HIV) virus, for which a comprehensive data set is available at the HIV-1, Human Protein Interaction Database http://www.ncbi.nlm.nih.gov/RefSeq/HIVInteractions. This data set collates interactions between the virus and human genes and proteins (protein-protein binding, effects on transcription, phosphorylation, expression, etc). Currently, 1510 viral-host interactions are reported from a total of 2977 known human proteins. The HIV-1 virus, thus, interacts in some way with about 5.2% of all known human proteins. Comparative figures as a percentage of 245 schizophrenia-related gene candidates are influenza = 21\%, HSV-1 = 22\%, CMV = 18\%, rubella = 12.6\%, and \textit{T. gondii} = 16\%, suggesting a generalized enrichment of pathogen-related genes in the schizophrenia gene data set. A more rigorous analysis is not currently possible given the incomplete data sets, both for host-pathogen interactions and for the growing list of genes implicated in schizophrenia.

In total, 114 of 245 susceptibility genes (46\%) can be related to the life cycles of one or other of these pathogens, a figure that is likely to increase when other species such as the Chlamydia family or the borna virus, also implicated as risk factors, are taken into account. These figures also compare favorably with the percentage of genes involved in endogenous brain processes related to the pathology of schizophrenia. For example, 24\% of the suspect genes encode for proteins related to the postsynaptic density, 16\% are involved in growth-related processes (eg, neuritic or axonal growth), 14\% can be related to oligodendrocyte function, 7\% to oxidative stress, and 7\% to dopaminergic function (see supplementary table 1 at http://www.polygenicpathways.co.uk/szfam.htm).

\textit{Susceptibility Genes Involved in Infection.} Human gene variants also modify infection, although there are relatively few genetic association studies in relation to the pathogens or genes reviewed in this study. Data from the Hugenet phenopedia\textsuperscript{131} (http://www.hugenavigator.net), which lists gene association studies for a number of infectious and other diseases, and a Medline search, are shown in supplementary table 2 (http://www.polygenicpathways.co.uk/infectassoc.htm), details of which are summarized in table 1. In all, 5 of 10 genes (50\%) (\textit{APOE}, \textit{HLA-B}, \textit{IL10}, \textit{MICB}, \textit{IL18RAP}) implicated in HSV-1 infection and 9 of 14 genes (61\%) (\textit{HLA-B}, \textit{HLA-DRB1}, \textit{IL1RN}, \textit{IL10}, \textit{IL12B}, \textit{IL18}, \textit{IL18R1}, \textit{IL18RAP}, TNF) implicated in CMV infection have also been implicated in schizophrenia. Although the numbers are small, and the genes involved almost exclusively related to the innate immune system, the enrichment does illustrate that certain genes implicated in schizophrenia are also related to infection. Clearly, it would be interesting to investigate the effects of polymorphisms in more relevant genes related to cerebral function such as \textit{NRG1} or \textit{DISC1}.

Of course, the physiological significance of any particular interaction relates to the role of the human gene/protein in the context of schizophrenia pathology, whether the
pathogen–host interaction is able to disrupt the role of the human gene in a relevant manner, or whether the host polymorphism is able to affect the pathogen life cycle. A number of whole-genome association analyses are in progress in relation to schizophrenia and related disorders. Antibody serotyping, pathogen DNA/RNA detection in the same blood samples, or a detailed knowledge of infection history would permit multifactorial analysis of any permutations of gene–pathogen–disease interactions. Such analyses are clearly warranted and would help to answer many of these questions. More direct analyses, e.g., viral/human 2-hybrid experiments using “normal” and variant susceptibility genes would also be informative.

Support From Microarray and Expression Studies. Indirect support for pathogen involvement, or immune activation in schizophrenia, derives from a series of microarray studies on schizophrenic brain tissue, which have shown marked changes in the expression of genes related to the immune and inflammatory system. In a collective analysis of multiple brain gene expression studies from the Stanley Research Consortium, genes related to multiple histocompatibility complex receptor activity and antigen processing and presentation figure among the top 10 of the Gene Ontology classified processes. In addition, a recent study has found that alpha-defensins are selectively elevated in T-cell lysates from schizophrenic patients. Defensins, which are produced by immunocompetent cells, and triggered by infection, are small endogenous peptide antibiotics with antiviral, bactericidal, and immunomodulatory properties. These studies clearly highlight the importance of pathogen and immune defense in relation to schizophrenia in adulthood.

Natural Selection and the Evolutionary Perspective. It has been suggested that schizophrenia susceptibility genes have been maintained in the human population at a frequency of about 1%, despite negative fitness, because of some form of positive selection pressure. In other words, certain gene variants must be of some benefit to the population. Many of these genes are implicated in pathogen life cycles and may well influence their virulence. For example, NRG1 signaling releases a factor that inhibits the replication of the influenza virus. Certain gene variants may favor the pathogens, e.g., polymorphisms in genes related to the IL18 signaling pathway or in the natural killer cell ligand MICB are associated with both herpes viral infection and with schizophrenia. Other variants associated with schizophrenia could well protect in some way against these agents. This is the case for a CCR5 delta32 deletion which protects against *T. gondii* and other infections. This deletion has also been implicated in schizophrenia but only in late-onset patients, suggesting either that it was associated with this particular type of disease or that it delays the onset of schizophrenia, a supposition that would be compatible with an anti-infective potential.

The mothers of schizophrenic patients born during influenza pandemics evidently survived the virus, but their progeny may bear the scars of the infection, e.g., activation of the eIF2alpha kinase signaling cascade by viral, cytokine, or stress-activated kinases. The selection pressure maintaining certain genes in the population could, thus, well be related to anti- as well as proviral effects, which would diminish mortality, at the expense of collateral damage, that nevertheless has devastating psychiatric consequences.

Genetic Implications

Association studies compare the frequencies of a particular gene variant between a normal and disease-affected population. They are notoriously inconsistent, and problems of replication between populations are the norm rather than the exception. Because these pathogens so clearly interact with many of the schizophrenia susceptibility genes during the course of their life cycles, many of the genetic associations could plausibly be related to extrinsic factors that contribute to the disease, as well as to the intrinsic disease process itself. It seems reasonable to assume that polymorphisms in such pathogen-related genes may well affect the infectivity or virulence of the pathogen, or the ability of the host to retaliate, and that the pathogens will in turn affect the function of the susceptibility genes, with which they interact.

Specific infections have distinct geographical and temporal distributions. If any particular virus interacts with a particular susceptibility gene or its product, the risk-promoting effects of the gene may be conditional on the presence of the virus at a particular time, when its effects are relevant to the genesis of particular subpathologies of schizophrenia (see also Pearce). In addition, genes and proteins are expressed at different times and in different tissues and cellular compartments during our lifespan, at times, which could coincide (or not) with infection or with the localization of the pathogen. Such conditional effects could well explain some of the heterogeneity in genetic studies. As shown in table 1, the relationship of different pathogens to different genes is similarly heterogeneous. It seems likely that a stratification of genetic association data in relation to pathogenic history could markedly influence the strength of association for many candidate genes.

Medical Implications and the Timing of Infection

A less than 100% concordance rate for schizophrenia in identical twins suggests that genes per se do not cause the disease. The pathogens implicated in schizophrenia are also common and are unlikely to be implicated in schizophrenia in all individuals. However, the convergence of genes and pathogens on a common signaling
cascade and a plethora of direct interactions between host and pathogen genes and proteins suggests how genetic variants could contribute to disease in infected individuals and how infections might contribute to disease in populations with particular genetic variants.

One could reasonably argue that, in some cases, these pathogens may well contribute to certain features of the disease but only in genetically susceptible individuals. If so, then elimination of the pathogens or prevention by vaccination could have a major effect on the incidence and perhaps the progression of the disease, as already suggested by several authors. \(^{7,143–145}\) Several infections implicated in schizophrenia occur many years before the onset of the disease, eg, maternal influenza infections during pregnancy or CMV or mumps infections during early childhood. There is no obvious indication that a small percentage of such patients might later develop a psychiatric disease and no reason why antiviral agents should be effective in later life when psychosis develops. However, vaccinations preventing such infections could have a significant effect on the incidence of psychiatric disease at later periods. A decline in the incidence of schizophrenia has, indeed, been reported in relation to the introduction of the polio vaccine. \(^{146}\) In addition, common childhood diseases, usually considered as benign, should perhaps be viewed from another perspective given the accumulating evidence that such infections can markedly increase the risk of developing psychiatric disorders later in life and a genetic rationale as to how this might affect particular individuals.

Other pathogens that have been detected in adulthood (eg, herpes viruses, Chlamydia, and \(T\) gondii) suggest that infections in adult life may also contribute to schizophrenia pathalogy. These may themselves be amenable to therapy, and their detection and elimination may have beneficial effects in schizophrenia. Interestingly, the use of valacyclovir has been reported to reduce symptomatology in schizophrenia patients seropositive for CMV. \(^{144}\) The use of activated immune cells coupled with antibiotic treatment has also been reported to improve psychiatric symptomatology in schizophrenic patients with Chlamydia infection. \(^{145}\) The situation may be complicated by the possibility that different and, perhaps, multiple pathogens may be implicated in different populations, although all converge on the same defence network (figure 4). Pathogen screening, perhaps, using antibody or gene arrays may well be useful in identifying these and other potential pathogens, some of which are amenable to therapy. Such therapy might, however, need to be tailored to individual pathogens.

**Conclusion**

Genes and pathogens both have a role to play in the genesis of schizophrenia. These factors appear to be interdependent. The pathways traced out by schizophrenia susceptibility genes relate both to intrinsic signaling networks and to the pathways specifically used by pathogens to commandeer host cell physiology. The host and pathogen genes and proteins together form an integrated transgenomic network (figure 4) that may govern the risk of developing schizophrenia. The genetic and medical implications are far reaching and likely to apply to the many other polygenic diseases in which pathogens also act as risk factors. Perhaps, an important take-home message is that many infections, generally considered as relatively benign, could in fact be a contributory factor to schizophrenia and that their identification, prevention, and treatment could have marked effects on the disease.

**Supplementary Material**

Supplementary tables 1 and 2 and other data are available at http://schizophreniabulletin.oxfordjournals.org.

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