Mutation in \( hprt1 \) Gene or HPRT Deficiency May be a Restricting Progeny of Favipiravir in Covid-19

Muhammad Torequl Islam

Department of Pharmacy, Life Science Faculty, Bangabandhu Sheikh Mujibur Rahman Science and Technology University, Gopalganj (Dhaka) - 8100, Bangladesh

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*Corresponding author: Muhammad Torequl Islam

Abstract

Favipiravir (FPV), a pyrazine carboxamide derivative (6-fluoro-3-hydroxy-2-pyrazinecarboxamide) is a broad-spectrum anti-viral drug which acts against many species of Arenaviridae, Bunyaviridae, Caliciviridae, Filoviridae, Flaviviridae, Orthomyxoviridae, Paramyxoviridae, Picornaviridae, Rhabdoviridae, Togaviridae groups. According to the WHO, FPV may be a new hope for the treatment severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) infections and for this it has undergone for a number of clinical trials. However, human hypoxanthine guanine phosphoribosyltransferase (HPRT) is believed to play a key role in its physiological activation process, therefore, the patients having Kelley-Seegmiller syndrome, Lesch–Nyhan syndrome, myocardial ischemia, hyperuricemia, anemia, and phosphoribosyl diphosphate (PRPP) synthetase superactivity due to deficiency this enzyme might be a limiting factor for the effectiveness of this drug in these types of patients.

Keywords: Favipiravir; SARS-CoV-2; Covid-19; Pathophysiology.

Favipiravir (FPV, also called avigan, favilavir, favipira, T-705), a pyrazinecarboxamide derivative was discovered by chemical modification of a pyrazine analog [1]. To date, it has been found to act against a number of viruses, including Orthomyxoviridae (Influenza: e.g., seasonal, H5N1, H1N1 pdm09, H7N9, A, B and C), Bunyaviridae (e.g., La Crosse, Punta Toro, Rift Valley fever, Sandfly fever, Dobrava, Maporal, Crimean-Congo hemorrhagic fever, Prospect Hill, and Severe fever thrombocytopenia syndrome), Arenaviridae (e.g., Junin, Pichinde, Tacaribe, Guanarito, Machupo, and Lassa), Filoviridae (e.g., Ebola), Rhabdoviridae (e.g., Rabies), Paramyxoviridae (e.g., Human metapneumovirus, Respiratory syncytial virus), Flaviviridae (e.g., West Nile, Yellow fever, and Zika virus), Togaviridae (e.g., Western equine encephalitis, Venezuelan equine encephalitis, Eastern equine encephalitis, Barmah Forest, Ross river, Mayaro, and Chikungunya), Picornaviridae (e.g., Polio, Rhino, and Enterovirus 71) and Caliciviridae (e.g., Noro) [2]. But, there is no record for its anti-coronavirus activity. Recent evidence suggests that this drug may act against life-threatening RNA virus infections [27]. And finally, it undergoes a number of clinical trials with the aim of coronavirus disease 2019 (Covid-19) treatment.

The exact mechanism of interaction of FPV ribofuranosyl-5′-triphosphate (FPV-RTP) with RNA-dependent RNA polymerase (RDpR) molecule is yet to be elucidated. It is hypothesized that FPV has virucidal effects, and it may be misincorporated in a nascent viral RNA, or it may act by binding to conserved polymerase domains, thus preventing incorporation of nucleotides for viral RNA replication and transcription [3]. It is demonstrated that FPV induced lethal mutagenic effect on influenza virus [1]. It is a prodrug that is metabolized to its active form, FPV-RTP [4]. Unlike an RNA virus, humans do not have RDpR, but have DNA-dependent RNA polymerase (DdRp) and DNA-dependent DNA polymerase (DdDp) [3].

In a study, FPV-RTP slightly inhibited human RNA polymerase II, which belongs to DdRp [4]. Human hypoxanthine guanine phosphoribosyltransferase (HPRT) is believed to play a key role in its activation process [4]. HPRT is an enzyme encoded in humans by the \( hprt1 \) gene, that catalyzes conversion of hypoxanthine to inosine monophosphate and guanine to guanosine monophosphate [5]. This reaction transfers the 5-phosphoribosyl group from 5-phosphoribosyl 1-pyrophosphate to the purine [6], therefore, it plays a key
role in the generation of purine nucleotides through the purine salvage pathway [7]. HPRT deficiency is inherited as a recessive X-linked trait [8], thus, males are generally affected and women are the asymptomatic carriers to its deficiency. Partial deficiency of the HPRT is also called Kelley-Seegmiller syndrome (a rare genetic disorder manifesting as a gout-uro lithiasis) [9].

The phosphoribosylpyrophosphate synthetase (PRPS) superactivity, a rare X-linked disorder resulting in increased activity of PRPS1, which is the gateway enzyme for the entrance of ribose into the purine pathway, catalyzing the conversion of adenosine triphosphate (ATP) and ribose-5-phosphate to 5-phospho-α-D-riboyl-1-pyrophosphate. PRPS1 superactivity results in increased purine production, leading to increased urate production and gout. It may also increase the function of mutated proteins [10]. HPRT deficiency and PRPS superactivity in cultured fibroblasts exhibited acceleration of purine synthesis de novo [11]. It seems HPRT deficiency may be linked to the PRPS superactivity.

In a study, HPRT enzyme has been seen to increase in non-small-cell lung cancer A549 and NCI-H460 cells [12]. Rosenstraus and Chasin [13] found that the genes responsible for glucose-6-phosphate dehydrogenase (G6PD) and HPRT activity are linked in Chinese hamster ovary cells. Pai et al., [14] also demonstrated a similar findings in human. G6PD deficiency occurs by g6pd gene mutation, is also inherited in an X-linked recessive manner and the most common medical problem include- hemolytic anemia, jaundice, dark urine, fatigue, shortness of breath, rapid heart rate, and so on [15].

Lesch–Nyhan syndrome (LNS) is a rare inherited disorder caused by a deficiency of the HPRT [16]. It is due to mutations in the hprt1 gene on the X chromosome [17]. The HPRT deficiency increases uric acid in our body fluids [18]. Generally, the combination of increased synthesis and decreased utilization of purines leads to high levels of uric acid production results severe gout and kidney problems, poor muscle control and moderate intellectual disability (e.g., self-mutilating behaviors, characterized by lip and finger biting), facial grimacing, involuntary writhing, and repetitive movements of the arms and legs similar to those seen in Huntington's disease [19]. It also causes neurologic and behavioral abnormalities, including mental retardation, spasticity, and choreoathetosis [18]. LNS may exhibit minor neurologic manifestations, including dysarthria, hypertreflexia, positive Babinski’s signs, or abnormalities of spinocerebellar function [20]. The cause of the neurological abnormalities may be linked to the poor utilization of vitamin B12 (some males may develop megaloblastic anemia) due to the deficiency of this enzyme [21]. The gene mutation in LNS occurs usually in the mother and passed on to her son (X-linked recessive manner). However, 33% of all cases arise de novo (from new mutations) and do not have a family history. It should be mentioned that the LNS is present at birth in baby boys (http://ghr.nlm.nih.gov/condition/lesch-nyhan-syndrome) [22]. Its deficiency has been found to link with myocardial ischemia [23] and hyperuricemia, juvenile-onset gouty arthritis, nephrolithiasis, and mild neurologic symptoms [24].

Moreover, the effectiveness of FPV against influenza pandemics is still controversial [25], as it may develop resistance against influenza virus [26]. Therefore, the effectiveness of this hopeful drug of the patients having a hprt1 gene mutation or deficiency of HPRT enzyme, especially who have Kelley-Seegmiller syndrome, Lesch–Nyhan syndrome, myocardial ischemia, hyperuricemia, anemia (e.g., hemolytic and megaloblastic), G6PD deficiency, cardiac diseases, and PRPS superactivity is an important issue. Moreover, FPV has broad-spectrum virucidal as well as genotoxic (on human RNA polymerase II which belongs to DdRp) and mutagenic (on viral cell) effects. Therefore, adequate research is urgently needed regarding the toxicogenetic profile (i.e., toxicity, cytotoxicity, genotoxicity and mutagenicity) to ensure the host genomic safety and stability during the long-term therapy with FPV.

Conflict of interest: None declared.

REFERENCES


