

An Insight to Herpes Zoster Review Article

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Abstract

Herpes Zoster (Or Simply Zoster) Is An Acute, Self- Limiting Viral Infection Characterized By Painful Vesicular Eruptions With Erythema Typically Present As Unilateral Dermatomal Rash. It Is Caused By Reactivation Of Dormant Varicella Zoster Virus. About 1 Million Patients Per Year Are Affected By This Condition. It Mainly Affects The Elderly And Persons With Waning Cell Mediated Immunity. If Left Untreated It May Lead To Various Complications Of Significant Morbidity Leaving A Considerable Effect On Quality Of Life As Well As Economic Status Of The Patient; The Most Serious Complication Being The Post Herpetic Neuralgia, A Chronic Neuropathic Pain Syndrome Which Leaves The Patient In A Debilitating State. This Review Article Provides An Overview Of The Disease And Emphasizes More On The Classical Features And Conventional Treatment Modalities Of Zoster Thus Enabling The Oral Physician To Make Early Diagnosis And Give Prompt Treatment, Which Is The Mainstay For The Management Of The Disease.

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20 **Index terms**— herpes zoster, shingles, zona, varicella zoster virus, zoster sine herpete, post herpetic neuralgia, 21 dermatomal rash, vaccine.

22 An Insight to Herpes Zoster Review Article Dr. V.Nagalaxmi ? , Dr. Anshul Singh ? , Dr. Srikanth K. ? , 23 Dr. Prameela K. ? & Dr. Swetha Reddy ¥ II.

1 Etiopathogenesis

25 HZ, with a lifetime risk of 10-30%, affecting about 1 million patients per year is caused by VZV. 1,4 VZV 26 belongs to alpha herpes virinae and consists of an icosahederal nucleocapsid enclosed in lipid envelope with 27 double stranded DNA at its centre. 2,3 The molecular weight and diameter is approximately 80 million and 28 150-200nm respectively. 2 HZ, a highly transmissible disease may spread either by respiratory droplets or direct 29 contact. 6 VZV first enters the host and causes infection of respiratory tract or epithelium of the conjunctiva. 6 It then replicates and

31 Author: e-mail: anshuls.4@gmail.com multiplies; and then penetrates the reticulo-endothelial system from 32 where the blood and lymphatics carry it throughout the body. 2,6 It then travels via mononuclear cells and 33 spreads to epidermis via capillary epithelium where VZV destroys basal cells. 6 This leads to generalized rash of 34 chickenpox. After the fall of the initial outbreak, VZV retreats into perineurial satellite cells of dorsal nerve root 35 ganglion where it remains inactive for years. 2,6 Reactivation of VZV by any triggering factor causes an outbreak 36 and the secondary infection of HZ. 2 Therefore, the primary infection by VZV causes chickenpox(varicella) 37 in children whereas shingles is caused by recurrent secondary infection in adults. 5 Incubation period for 38 varicella ranges from 14-16 days; chances of transmission being high between 10-21 days after initial exposure. 39 2 Transmission cannot occur after crust have dried. Indirect transmission does not occur. 2 Most commonly 40 affected dermatomes are thoracic (45%), cervical (23%) & trigeminal (15%). 7 HZ may affect sensory ganglia & 41 its cutaneous nerves (Strommen et al. 1988) 6 . Thoracic and lumbar dermatomes are involved more commonly 42 as compared to craniofacial area. 8 The virus may remain latent for decades together in the cranial nerve, 43 dorsal root and autonomous nervous system ganglia along the entire neural axis. 9 2 main mechanisms have

4 CLINICAL FEATURES

44 been developed by VZV to escape the human immune system: 3 a. Initially, VZV remains inactive in sensory
45 ganglion, thereby restricts the expression of viral proteins. At this stage virus does not replicate but retains its
46 capability to revert to pathogenic nature at anytime. 3 b. Down regulating the expression of antigens of MHC
47 Class 1 on the surface of infected cells, leads to decrease in surface expression of its proteins, thereby restricts
48 the presentation of vital peptides to cytotoxic T-cells which ultimately leads to escape of lysis by virus infected
49 cells. 3 Most critical complication is a form of neuropathy of pain called post herpetic neuralgia (PHN). 9 The
50 pathophysiology involved is injury affecting the neurons of both central and peripheral nervous system generates
51 spontaneous discharges. 4 It also decreases the action potential threshold which in turn decreases the generation
52 of disproportionate pain, even with non-specific stimuli. 4

53 2 III. Epidemiology & Predisposing Factors

54 HZ, a common disease with a lifetime risk of 10-30% which increases to 50% among individuals limiting viral
55 infection characterized by painful vesicular eruptions with erythema typically present as unilateral dermatomal
56 rash. It is caused by reactivation of dormant varicella zoster virus. About 1 million patients per year are affected
57 by this condition. It mainly affects the elderly and persons with waning cell mediated immunity. If left untreated
58 it may lead to various complications of significant morbidity leaving a considerable effect on quality of life as
59 well as economic status of the patient; the most serious complication being the post herpetic neuralgia, a chronic
60 neuropathic pain syndrome which leaves the patient in a debilitating state. This review article provides an
61 overview of the disease and emphasizes more on the classical features and conventional treatment modalities of
62 zoster thus enabling the oral physician to make early diagnosis and give prompt treatment, which is the mainstay
63 for the management of the disease.

64 3 Introduction

65 herpes zoster also called zona or shingles is a common viral disease among the elderly and immunocompromised,
66 is unilateral and associated with painful vesicular dermatomal skin rash and vesicles, frequently in a striped
67 pattern. 1 Reactivation of varicella zoster virus (vzv) causes herpes zoster (hz). 2 HZ is derived from greek
68 word herpein meaning to creep or spread; zoster meaning girdle or zone, hence the name zona (warrior armour
69 binding in a belt-like fashion). 2,3 Shingles, derived from latin cingulum meaning girdle (unilateral rash that
70 enfolds like a girdle around the torso). 3 H ?85years. 1 In Australia by the age of 30 years more than 97%
71 of population have antibodies to VZV, which confirms that they have been already infected with virus. Thus,
72 the entire adult population is at a high risk of HZ. 5 1.2-4.8 per 1000 people per year is the total incidence
73 among immunocompetent persons. 1 HZ ranges from 14.5-53.6 per 1000 persons-years in immunosuppressed
74 patients. 1 HZ increases with age with approximately 14.2 per 1000 people per year in persons ?50 years in
75 USA,UK,Italy and Germany. 1 Recurrence is seen in approximately 4 % of patients who develop HZ. 10 HIV
76 patients are 10 times more prone to develop HZ compared to general population. 1 HZ in organ transplant
77 patients ranges from approximately 22-per 1000 personyears overall, with increased predilection among African-
78 American patients (37.6 per 1000 persons-years) and heart transplant patients(40 per 1000 person-years). 1 HZ
79 incidence is increased in patients treated with mononuclear antibody-TNF inhibitors and various biologics (19.1
80 per 1000-person-years) compared to non-systemic therapy patients (4.6 per 1000 personyears). 1 HZ is more
81 liable in individuals who suffer with leukemia, lymphoma, metastatic malignancy, autoimmune disorders like
82 SLE, RA, Wegener's Granulomatosis, Diabetes, COPD, Patients on cytotoxic drugs or steroids & those receiving
83 chemotherapy. 1 Psychological stress may also contribute to HZ. 3 Female predilection for HZ (Thomas and Hall's
84 systematic review). 1 Malnourishment leads to decrease in cellmediated immunity thus increases susceptibility to
85 HZ. Alcohol and smoking affect on HZ is still unclear. 1 Climatic changes also influences shingles wherein persons
86 residing in temperate climate and northern latitude are at an increased rate of developing shingles. 3 Another
87 risk factor to HZ is mechanical trauma and immunotoxin exposure. 1 Prior infection with VZV (chickenpox,
88 vaccine) is an important predisposing factor for the development of HZ. 5 Association between varicella & HZ
89 was first made in 1892. 11 IV.

90 4 Clinical Features

91 HZ presents as an acute, sporadic, self-limiting, painful unilateral vesicular dermatomal rash, often lasts for
92 approximately 10-15 days. 5 Pain and rash are the cardinal features of HZ. 12 The prodromal (pre-eruptive)
93 stage is characterized by pain which may be intermittent/continuous, boring, tingling, itching, burning, prickling
94 or knife-like in the epithelium surface supplied by the affected sensory nerve. 3,13 This severe neuralgia is caused
95 due to viral replication which in turn leads to active ganglionitis with resultant neuronal necrosis. 13 Prodrome
96 may also be associated with mild fever, headache, malaise, dysesthesia. 3 The cutaneous features are preceded
97 by prodromal stage (continue for 3-5 days) in 80% patients (Strommen et al. 1988, Carmichael 1991, Millar
98 & Troulis 1994). 3,6 Odontalgia may be the only oral manifestation present at this stage (Barrett et al. 1993,
99 Law & Lilly 1995). 6 The Acute (active) phase is characterised by unilateral dermatomal rash associated with
100 malaise, headache, mild fever and nausea. Rash appears proximally and spreads distally. 3 The rash advances in
101 12-24 hours from erythematous papules and oedema to vesicles and finally within 1-7 days it advances to form
102 pustules. 6 The pustules then dry and form painful crust which within 14-21 days fall-off, therefore leading to

103 formation of macular and erythematous lesions which usually heals to form hypo/hyper pigmented scars. 6 In
104 severe cases, hemorrhagic necrosis may lead to loss of areas of epidermis and dermis (Strommen et al. 1988,
105 Carmichael 1991). 6 Intraoral lesions usually appear after cutaneous rash. 3 HZ without rash, condition termed
106 as Zoster sine herpete, is seen in rare cases wherein the affected patients suffer with pain which is sudden, severe
107 and hyperesthesia over a specific dermatome. 13 Chronic neuropathic pain syndrome stage is also called as Post
108 Herpetic Neuralgia (PHN). 6 Dworkin defined PHN as "a significant pain or abnormal sensation 120 days or more
109 after the presence of initial rash." 4 It described as pain comprising of 3 prominent components: 6 I. Constant,
110 usually deep pain II.

111 Brief, recurrent shooting pain III.

112 Allodynia -sharp, radiating dysesthetic sensation caused by even slight touching (Rowbotham & Fields 1989).
113 V.

114 **5 Oral Manifestations**

115 Oral complications are seen when HZ affects the Trigeminal Nerve (18-20% cases). 11 Unilateral multiple
116 vesicular eruptions (1-4 mm) with erythema is seen intra orally. 11,13 Vesicles on palate, uvula, tonsils, tongue,
117 buccal mucosa and floor of the mouth are seen depending upon the branch involved. 11 Apart from odontalgia,
118 devitalised teeth, internal resorption, pulpal necrosis, developmental anomalies, sudden exfoliation of teeth, facial
119 scarring, jaw osteonecrosis, severe periodontitis may also be appreciated.

120 **6 Investigations and Diagnosis**

121 Pain, Unilateral nature and Segmental distribution accounts for clinical diagnosis of HZ. 3 Laboratory
122 tests include Tzanck Smear, Viral culture (30-70% sensitive; 100% specific), FNAC from fresh vesicles. 3
123 Molecular techniques such as Dot-Blot hybridization and Polymerase Chain Reaction for detection of VZV DNA
124 (approximately 100% sensitive). 11,13 Direct Immunofluorescence assay is a good diagnostic aid. 11 VIII.

125 **7 Differential Diagnosis**

126 Differential Diagnosis may include Trigeminal neuralgia, Maxillary sinusitis, Periodic Migranous neuralgia,
127 Myocardial pain, Atypical facial pain, Munchausen's Syndrome(Drinnan 1987). 6 The Prodromal stage pain
128 can be misdiagnosed as Pleurisy, Thrombophlebitis, Cardiac disease, Duodenal ulcer, Cholecystitis, Bell's Palsy,
129 Otitis media, Herniated nucleus pulposus, Sensitive teeth. 11,13 IX.

130 **8 Management**

131 The primary management comprises of early diagnosis and prompt treatment in the prodromal stage. Management
132 is emphasized towards pain control along with prevention of PHN, supportive care and hydration and
133 definite treatment to decrease the dissemination may be isolated to avoid cross-infection and complete bed rest
134 may be advised. Hospitalization is advised for immunocompromised patients.

135 Antiviral drugs have been proven to decrease the pain and duration of rash, as well as speed up healing and
136 prevent further complications. 3 Care should be taken to administer antivirals within 72 hours after onset of rash.
137 Prednisolone (60 mg daily initially, care should be taken to taper the dose for 21 days) may be useful in reducing
138 acute pain. 12 Some cases have been treated with Amitriptyline 25 mg/day for 3 months to prevent PHN. 12 Relief
139 from severe acute pain by administering single epidural injection of corticosteroids (80 mg methylprednisolone)
140 and Local anesthetic (10 mg bupivacaine) may be effective. 16 Opioids and NSAID's has been proven to be
141 effective to relieve acute pain. Oxycodone decreases acute pain and tramadol prevents PHN. 12 XI.

142 **9 Treatment for Post Herpetic Neuralgia**

143 The main objective of PHN treatment is to relieve pain and require a diverse approach. Multiple medications may
144 be needed. 3 The first line of treatment for PHN comprises of anticonvulsants like Phenytoin / Carbamazepine /
145 Gabapentin (100-300 mg/day orally at bedtime). Dosage may be increased until therapy is effective and response
146 appreciated but one should be cautious and should keep a constant check on the blood drug level. 11 Topical
147 application of 80% capsaicin cream (3-5 times daily) and 5 % lidocaine patch (every 4-12 hours) and Aspirin cream.
148 8,11,12 The second line of treatment is with opioid analgesics and tricyclic antidepressants like Amitriptyline
149 / Desipramine / Imipramine / Nortryptiline (25 mg/day orally at bedtime). Dosage can be increased until
150 sufficient response is met but maximum dosage should not exceed 150 mg/day. 8,11 Systemic Corticosteroids to
151 prevent PHN is controversial. Combination of intralesional steroids and Local anesthetic's have been proposed
152 to hasten healing and prevent PHN. 11 Selective Serotonin Norepinephrine Reuptake Inhibitors (SSRI's) may
153 be administered in patients who cannot tolerate TCA's. 3 Newer advances: 14 Argyll-Robertson pupil signifies
154 involvement of ciliary ganglia. 14,15 Ramsay Hunt Syndrome (triad of HZ of external ear, auditory symptoms,
155 ipsilateral facial paralysis) signifies involvement of geniculate ganglion. 11 risk especially in immunosuppressed
156 patients. 8 Patient -Spinal cord stimulation -Botulinum toxin injection X. Treatment for Herpes Zoster

157 In 1995, Varicella vaccine was recommended in USA for healthy children >1 year old, susceptible adolescents
158 and also adults. 1 In 2006, the FDA recommended a live attenuated vaccine derived from the oka strain of VZV

10 CONCLUSION

159 for prevention of HZ and its complications. 12 Since then a decrease of 90-95% of VZV infection in children aged
160 1-9 years was observed. 4 It is safe, well-tolerated cost-effective and efficient. Protection by the vaccine remains
161 for about 7 years. ??9 A single 0.65 ml dose injected subcutaneously in the deltoid region. ??0 Vaccine cause an
162 upgrade in cellmediated immunity thereby causing a decrease in shingles and also decreased incidence of PHN.
163 4 It also decreases the burden of illness. 5 Studies have shown a decrease of 51.3% in incidence of HZ; 66.5% in
164 incidence of PHN; 61% in BOI score. 5 FDA recommends HZ vaccine for adults ? 50 years irrespective of person
165 suffering with prior HZ episode. 3 ACIP (Advisory Committee on Immunization Practices) has not applied any
166 upper age limit for vaccine. 3 Care should be taken to increase the vaccination coverage if zoster vaccine is given
167 simultaneously with other vaccine. 1 Several studies are being conducted on effects of inactivated live attenuated
168 vaccine is not recommended. 3 Vaccine should be kept frozen at -15?C(once opened should be used within half
169 an hour). 3 FDA has approved transportation and storage at 2-8?C and upto 72 hours. ??0 Contraindications
170 may include cases of life threatening hypersensitivity reactions, HIV patients with CD 4 count <200, Patients on
171 chemo/radiotherapy, Pregnancy and Breast feeding. 3 XIII.

172 10 Conclusion

173 HZ though being a self-limiting condition, if left untreated can lead to various complications involving almost
174 all the organs of human system, with PHN being the most critical one. However, an oral physician can be the
175 first one to recognize the signs and symptoms thereby, being the first ones to make the initial diagnosis. Thus,
176 dentists should have complete knowledge about the disease so that prompt treatment can be given and patient
management can be done early and efficiently.



Figure 1:

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