

Primary Cutaneous *Adiaspiromycosis*: A Case Report and Review of the Literature

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ABSTRACT

Aim: To document a case of primary cutaneous *Adiaspiromycosis*.

Introduction: *Adiaspiromycosis* is a rare emerging fungal infection caused by *Chrysosporium Parvum* var *Crescens* now called as “*Emmonsia Crescens*” belonging to the genus *Emmonsia*, order *Onygenales*, family *Ajellomycetes*.

Case Report: A 38-year-old male presented with an ulcerated nodule over the infra-axillary region with itching for 3 months. The lesion began as small nodule slowly growing to attain the present size. His routine investigations were within normal limits. Serological tests such as HBsAg and HIV were nonreactive. Fine needle aspiration cytology showed ‘suppurative granulomatous’ inflammation. The nodule was excised and sent for histopathological examination. Histopathology showed numerous noncaseating granulomas and suppuration. Amidst suppuration, as well as inside the giant cells and within granulomas, numerous varying sized, thick-walled, round to oval ‘adialspores’ were seen. There was no evidence of budding or septation or endosporulation. They were PAS and GMS positive. He was treated with topical luliconazole and oral fluconazole. There was no recurrence on follow up of one years.

Conclusion: *Adiaspiromycosis* is a rare fungal infection and primary cutaneous involvement is a rare distinct entity. Detailed morphological assessment in histopathology is essential for its identification as the organisms are difficult to isolate in fungal culture from human clinical material.

Keywords: *Adiaspiromycosis*, *Haplomycosis*, *Chrysosporium parvum* var *crescens*, *Emmonsia crescens*, *Adiaspirosis*

INTRODUCTION

Adiaspiromycosis (or *Adiaspirosis* or *Haplomycosis*) is a rare emerging fungal infection caused by *Chrysosporium Parvum* var *Crescens* (*Emmonsia Crescens*). The disease is more common amongst primates and domestic animals and is rare in humans. Humans are accidental hosts and dead end of life cycle for the fungi. Humans contract the infection either by inhalation or inoculation of ‘spores.’ The lungs are the most common site affected but other sites can be involved. They usually remain localized to the site of entry, which only grow in size without actual division, to form large sized spores called ‘adialspores’ that elicit suppurative granulomatous inflammation and form a space-occupying lesion called ‘*Adiaspiroma*’. Dissemination is rare. Histopathology is the mainstay in diagnosis. Often the lesion is an incidental finding during histopathological examination of tissue. The organisms are difficult to isolate from human clinical material (1-5).

Primary cutaneous involvement is a rare distinctive entity. Here we are presenting a case report of cutaneous involvement of *adialspiro*mycosis.

CASE HISTORY

A 38-year-old male timber yard worker presented with an ulcerated nodule over the infraaxillary region with itching for 3 months. The lesion began as small nodule slowly growing to attain the present size. There was no history of fever, trauma, diabetes and hypertension or tuberculosis. On examination, there was a subcutaneous ulcerated nodule over the left infraaxillary region measuring 2 cm x 1 cm x 0.8 cm, freely mobile and slightly tender. His cardiovascular, respiratory and central nervous system functions appeared normal. His chest x ray was unremarkable. His routine investigations revealed hemoglobin of 13.88 gm%, total leukocyte count of $8.340 \times 10^3/\mu\text{L}$, platelet count of $320 \times 10^3/\mu\text{L}$, and the differential count showed 49% polymorphs, 45% lymphocytes, 5% eosinophils, and 1% mono-

cyte. His random blood sugar was 118 mg% and serological tests such as HBsAg and HIV were nonreactive. Fine needle aspiration cytology showed a ‘suppurative granulomatous’ inflammation. The nodule was excised and sent for histopathological examination.

Routinely processed Hematoxylin & Eosin-stained paraffin sections showed numerous discrete nodular areas interspersed with suppuration and fibrocollagenous and fibroadipose tissue. Nodular areas showed discrete well-delineated epithelioid cell granulomas with both foreign body and Langhans type giant cells. Most granulomas were nonnecrotic; however some were disrupted by central suppuration and some coalesced to form larger granulomas (Figure 1A).

Amidst suppuration as well as within the granulomas and giant cells were seen ‘round to oval ‘unicellular structures (Adiaspores) ranging from 25 to 80µm diameter with a thick trilaminated wall ranging from 6- 20 µm wide (Figure 1, 2). The laminated wall showed three distinctive zones

with the outer zone densely pink, wider middle zone faintly amphophilic, and inner zone faintly pink (Figure 2B). The majority of these spores were ‘empty’ looking rings. Some of them showed basophilic to amphophilic granular material diffusely as well as in aggregates abutting the inner wall (Figure 1B). Some were degenerated, collapsed, either ingested by multinucleated giant cells or lying freely amidst suppuration or giant cells seen entering into the fragmented large spores (Figure 3F, 4G). Some of the spores showed eosinophilic ‘Splendore Hoespli’ material around them with adherent neutrophils (Figure 1E). Suppuration showed neutrophils, degenerated nuclear debris, macrophages, lymphocytes, plasma cells and some eosinophils (Figure 1,2). Some larger irregular ovoid spores looked like broad aseptate non-branching hyphae (Figure 2C, D). Some of the old degenerating spores showed retraction of the inner wall simulating septation (‘pseudo-septation’) (Figure 1C, 2D, 3B, 4E). There was no evidence of ‘budding’ or ‘endosporulation’ or typical ‘septation’.

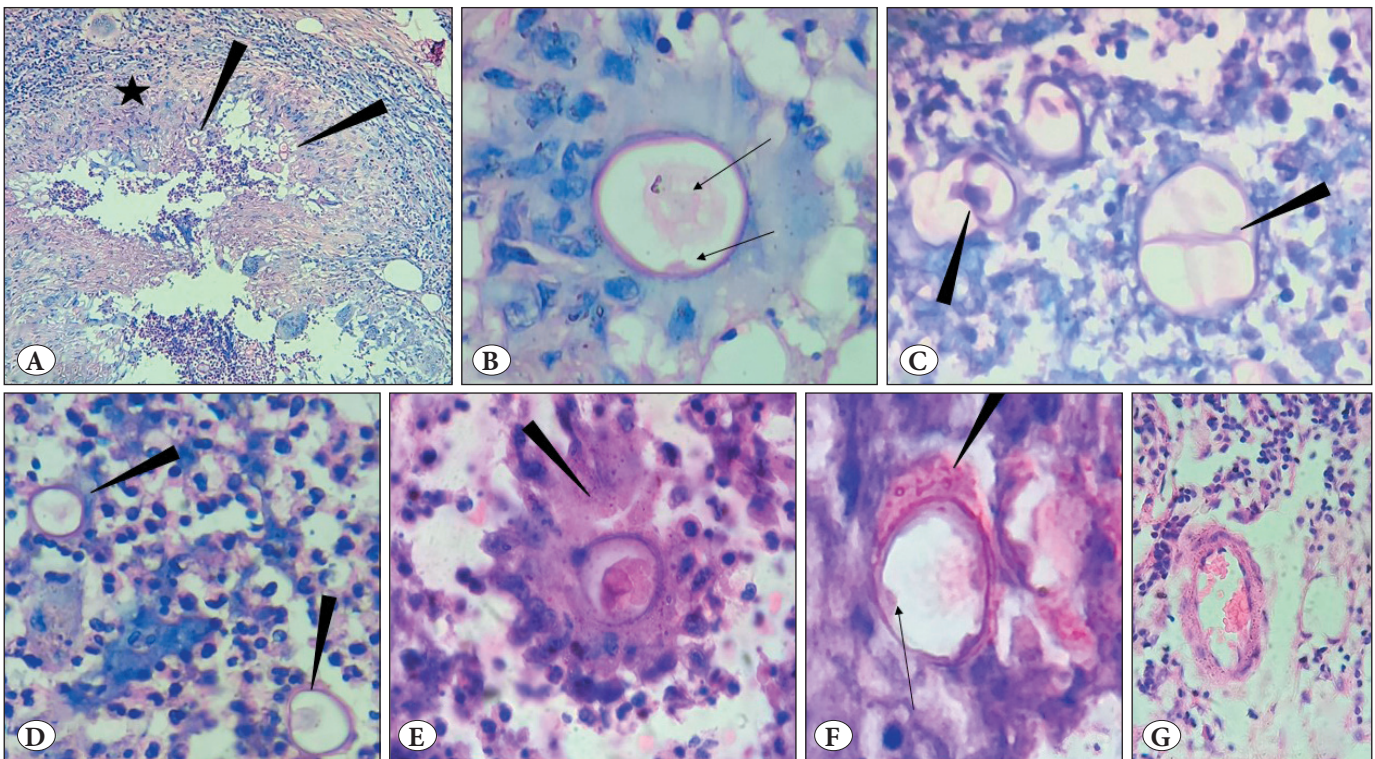


Figure 1: Showing ‘adidaspores’ with suppurative and granulomatous inflammation. **A)** Well-formed epithelioid cell granulomas (Asterix) with central suppuration and ‘adidaspores’ (black arrow heads) (x 10, H&E), **B)** large ‘adidaspore’ within multinucleated giant cell, with amphophilic to eosinophilic amorphous granular contents in the centre and eosinophilic granular content adherent to inner wall (black arrows) (x40, H&E), **C)** retraction of inner wall simulating septation (pseudo septation) (black arrow heads) (x100, H&E), **D)** Thick walled ‘adidaspores’ amidst suppuration (black arrow heads) (x40, H&E), **E)** Thick walled ‘adidaspore’ surrounded by ‘Splendore Hoespli’ material with a collar of adherent neutrophils (black arrow head)(x 100, H&E), **F)** tangential section through ‘adidaspore’ creating thick wall on one side (black arrow head) with granular contents on inner wall (black arrow) (x100, H&E), **G)** capillary with definite endothelial lining simulating ‘adidaspore’ (x100, H&E)

Special stain periodic Acid Schiff stain showed well-delineated spores with refractile pink trilaminated thick wall with adherent aggregates of pink granular material (Figure 3B, C).

Special stain Gomori's Methenamine Silver showed well-delineated large spores with brown black thick laminated wall and empty cytoplasm. Some showed faint gray granular material inside the cytoplasm (Figure 4).

Culture was not possible as there was no clinical suspicion and hence tissue received was formalin fixed. The case was diagnosed based on histomorphology as 'Adiaspiromycosis'.

He was treated with topical Luliconazole for two weeks and oral Fluconazole for 7 days. There was complete healing of the site with no recurrence on follow up for one year.

DISCUSSION

Adiaspiromycosis is a chronic fungal infection caused by the dimorphic fungi *Chrysosporium Parvum* var *Crescens* now called "*Emmonsia Crescens*" belonging to the genus *Emmonsia*, order Onygenales, family Ajellomycetes. Other members of the family Ajellomycetes include *Blastomyces*, *Histoplasma*, *Paracoccidioidomyces*, *Emergomyces* and

Emmonsiiellosis. Genus *Emmonsia* includes *E crescens*, *E parva*, *E pasteuriana* and *E Africanus*. Only *E crescens* and *E parvum* produce 'adidaspores' while *E pasteuriana* and *E Africanus* produce small budding yeasts but not adidaspores. *E crescens* produces larger 'adidaspores' and are considered responsible for "Adiaspiromycosis", while *E parva* produces smaller adidaspores and are reclassified as *Blastomyces parva* (*Blastomyces* Spp) (1-6).

The disease is relatively rare in humans where they are accidental hosts. The disease is prevalent amongst rodents and domestic animals (7-10). Humans contract the infection by inhalation or inoculation of spores. Most of the documented cases are localized at the site of entry with the lung being the most involved site (11-14). Rarely the spore disseminates to distant sites, particularly in immunocompromised hosts (15). Other sites of involvement include the bone, nose, paranasal sinuses, eye and heart (16). Primary cutaneous involvement is a rare distinct entity (16-21).

The organisms are soil saprophytes, and exhibit thermal dimorphism with asexual reproduction often occurring at room temperature where hyphal forms produce 'conidia' often directly from hyphae ('lollipop' like conidia also

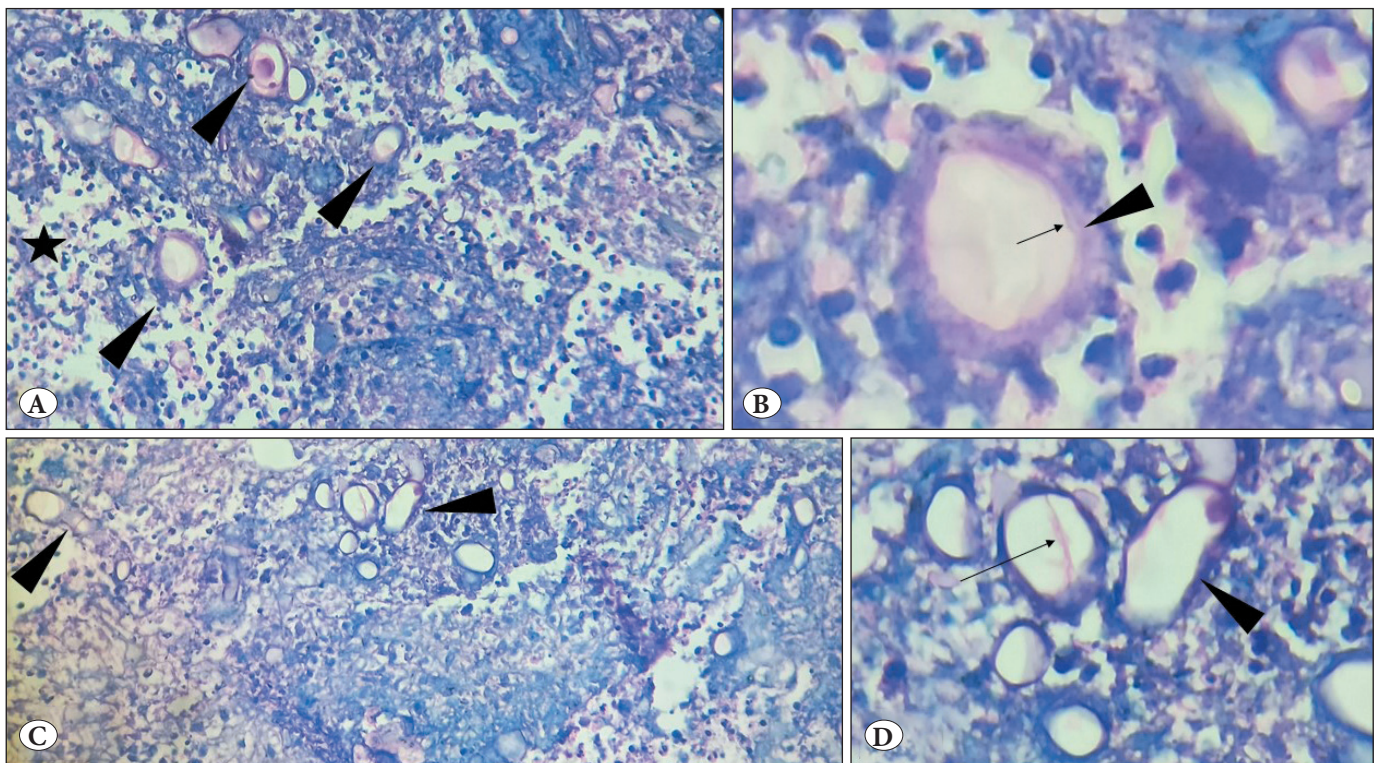


Figure 2: A) Showing numerous thick walled adiaspores (black arrow heads) amidst suppuration (Asterix) (x 40, H&E), B) Inset of A: trilaminar thick wall of adiaspore showing outer thick eosinophilic zone (black long arrow head), wider pale pink middle zone and inner pale irregular zone (black arrow) (x200, H&E). C) elongated adiaspore simulating 'hyphae' (black arrow heads) (x100, H&E), D) inset of C. adiaspore showing retraction of inner wall (black arrow) (x200, H&E).

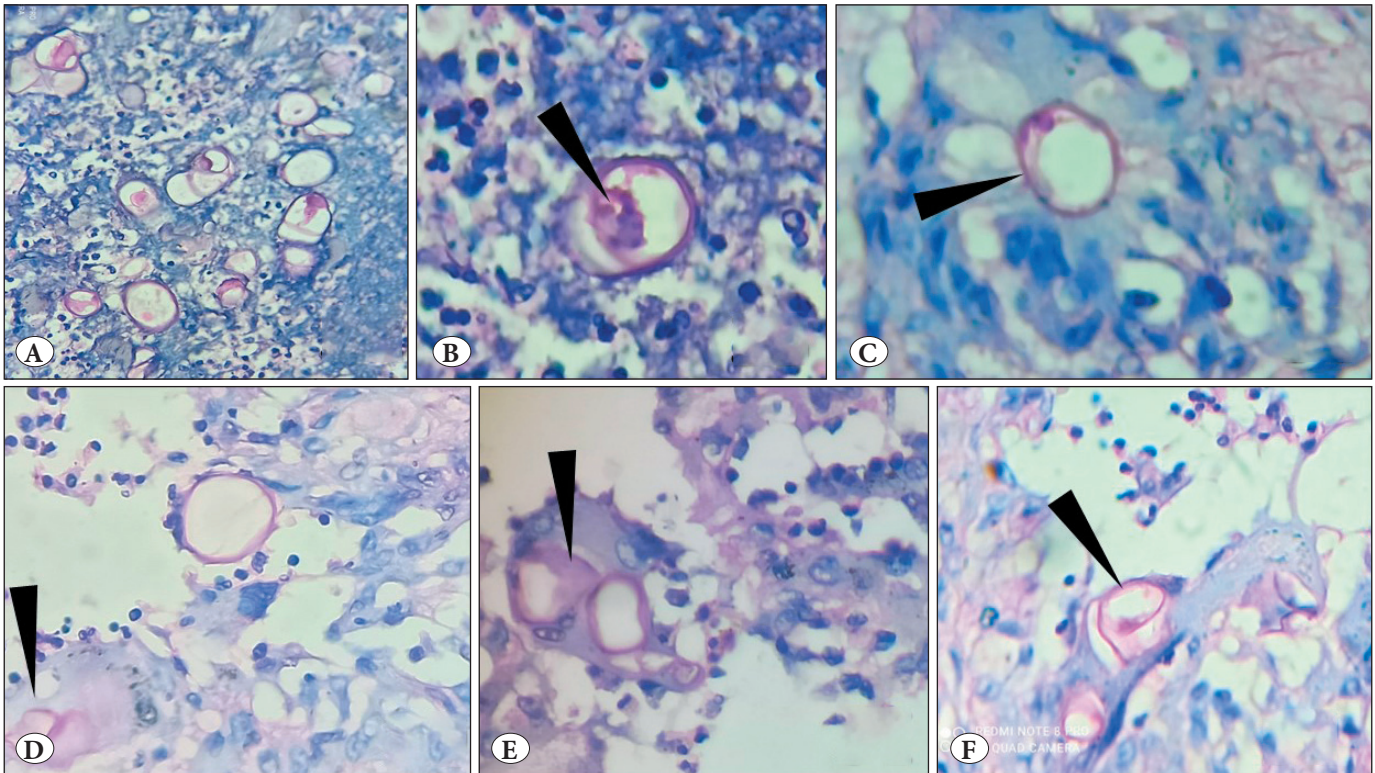


Figure 3: Showing Periodic acid Schiff positive ‘adiaspores.’ **A)** ‘adiaspores’ amidst suppuration (x40, PAS), **B)** adiaspore with retraction of inner wall on one side (black arrow head) with adherent irregular amorphous greyish pink granular material (x100, PAS), **C)** ‘adiaspore’ amidst granuloma (black arrow head) (x100, PAS), **D,E)** ‘adiaspore’ inside multinucleated giant cell (black arrow heads) (x40, PAS), **F)** degenerating & collapsed adiaspore inside a multinucleated giant cell (black arrow head) (x 40, PAS).

called ‘*aleurioconidia*’). The conidia that are released into the external environment grow and produce hyphae and the cycle continues (3,4,19).

In human tissue, exclusively only spores (‘conidia’) are seen and mycelial forms are rarely seen. They grow in size without multiplication, a unique feature amongst fungi (hence ‘non-replicating’ spores also called ‘Haplospores’ and the disease “*Haplomycosis*”). With time they enlarge to form large ‘adiaspores’, and elicit a granulomatous response with fibrosis forming characteristic ‘*adiaspiroma*’. They do not produce hyphae, and hence it is uncommon to see hyphal structures in tissue (1-4,11,15). In human tissue (37°C), ‘adiaspores’ lose their biological activity and viability. They enlarge in the direction of less tissue resistance and hence tend to take various shapes including ovoid, ellipsoid, cylindrical, and large ‘septate non branching hyphae like’ (Figure 2C, D). They develop distinctive refractile thick walls (20- 30µm wide) that are bilaminar or trilaminar. In Hematoxylin & Eosin-stained sections, the thick chitinous outer wall appears eosinophilic to amphophilic, while the inner wall is irregular and stains faintly eosinophilic. Both PAS and GMS delineate the laminated wall into three

distinctive zones as an intensely dark staining outer wall, wider faintly stained middle zone, and moderately stained wider irregular inner wall (1-3).

With time, the nuclei undergo degeneration and disappear or form precipitate or aggregates, and hence fully developed ‘adiaspores’ are ‘empty looking.’ In H&E-stained sections, the cytoplasmic contents including precipitated nuclear material appear as unstained or faintly stained amorphous granules in the center or may get aggregated at the inner portion of the thick wall adding to the substance of the thick wall. They appear faintly pinkish in PAS stain and faint gray in GMS stain. Sometimes the inner wall retracts from the outer wall creating a large empty space that may look like ‘central septation’ (pseudo-septation) and is likely to be confused with division of the organism (Figure 1C,2D, 4E). The structure of the inner wall is relatively complex, exhibiting irregular projections (Figure 2) and are ‘porous’ in nature simulating ‘honey combing’ or ‘mesh like’ ultra-structurally. The other differential stain to characterize the distinct zones include Alcian Blue which typically stains the inner wall red and outer wall blue (11).

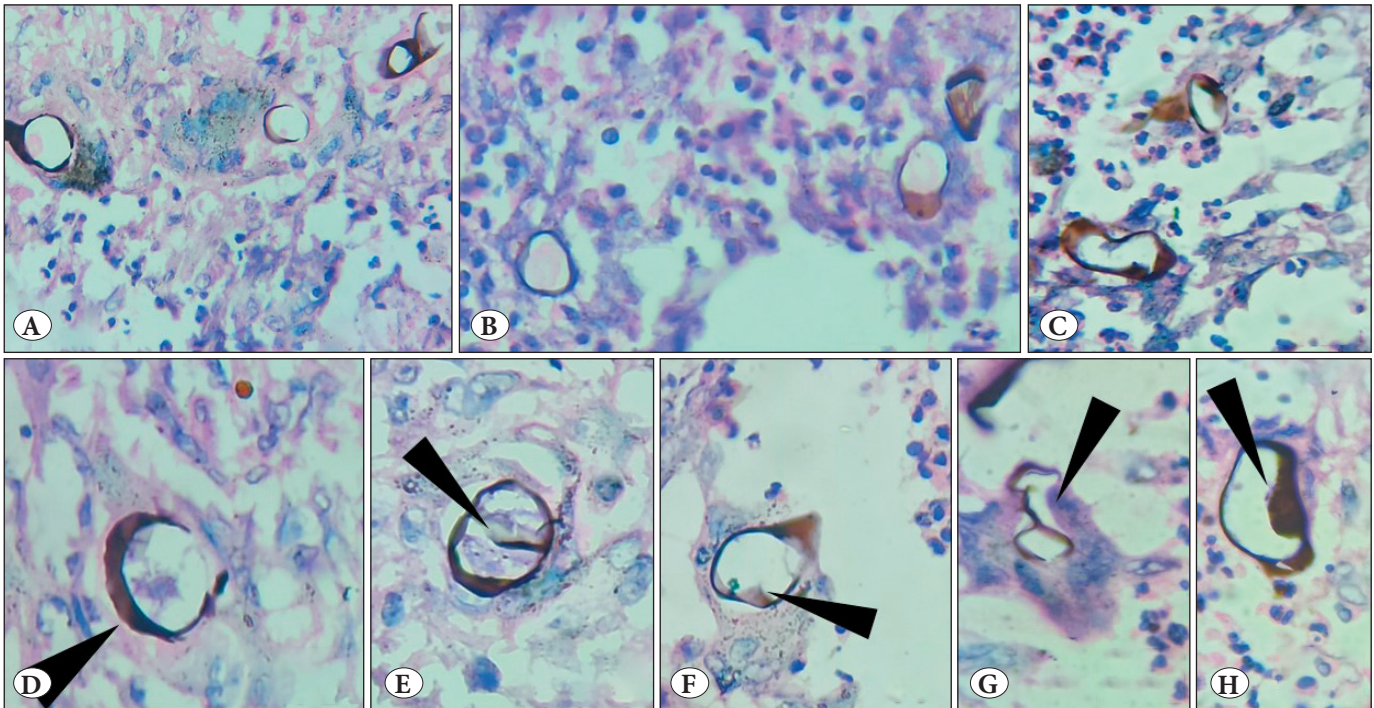


Figure 4: Showing Gomori's methenamine silver stain positive adiaspores. **A)** varying sized 'adiaspores' amidst granulomas (x40, GMS-HE), **B,C)** 'adiaspores' amidst suppuration (x40, GMS-HE), **D)** 'adiaspore' inside a multinucleated giant cell showing thick wall (black arrow head) with irregular greyish amorphous granular material inside (x100, GMS-HE), **E)** 'adiaspore' showing retraction of inner wall simulating septation (pseudo septation) (black arrow head) (x100, GMS-HE), **F)** 'adiaspore' showing greyish granular contents adherent to inner wall (black arrow head) (x100, GMS-HE), **G,H)** 'adiaspore' showing degeneration and collapse inside multinucleated giant cell (black arrow heads) (x100, GMS-HE).

They elicit a suppurative and granulomatous response with typical epithelioid cells, lymphocytes, and multinucleate giant cells engulfing empty rings of 'adiaspores' that may collapse or rupture. These empty rings of 'adiaspores' are seen amidst suppuration where they are surrounded by a collar of neutrophils (Figure 3F) or may exhibit an antigen antibody reaction with deposition of 'Splendore Hoespli' material (Figure 1F). Characteristically, necrosis and calcification are absent. Typically, the granulomas are 'non-necrotic', and hence calcification is rare and are walled by fibrous tissue, forming characteristic '*Adiaspiroma*' (1-4,14).

The size of the 'adiaspore' will give a clue to the duration of lesion. It may reach sizes of up to 50- 80 μm or more. On average it takes about 6- 8 weeks for spore to reach the size of 100 μm and it may reach a size up to 500 μm in long standing cases. Enlarging adiaspores may compress smaller airways with respiratory compromise (1,11,20,21).

As the lesion gets older, adiaspores tend to degenerate, become brittle and easily fold, fragment, become granular, and some become firm to hard so that during microtomy the entire spore gets dislodged by the microtome knife and hence one may see central empty space in granulomas (1).

Diagnosis

Histopathology is the mainstay in diagnosis. Most of the cases are diagnosed as an incidental finding (1-3,13). Culture of the organism is only possible from environmental sources and infected animals but not from human clinical material due to non-viability of 'adiaspores'. It is possible that human body temperature of 37°C does not favor not only the reproductive process but also the viability of the organisms. The fact that *adiaspores* are trapped inside thick-walled granulomas may make them nonviable (13). Hence, human tissue is the dead end of the life cycle for the fungi. The organism is difficult to isolate in culture from human clinical material. The diagnosis of *Adiaspiromycosis* mainly depends on their characteristic morphology in histopathology (1-4,13,14). There are no well-established skin tests and serological tests to diagnose the disease (3).

Differential diagnosis of *Adiaspiromycosis* include *Blastomycosis*, *Rhinosporidium Seeberi*, *Coccidioidomycosis*, *Mucorales*, helminths such as *Dirofilaria* & *Strongyloides* and small blood vessels (Table I).

The empty rings of 'adiaspores' are confused for empty spherules of *coccidioidomycosis* & *Rhinosporidium seeberi*,

Table I: Differential diagnosis of *Adiaspiromycosis*.

<i>Blastomycosis</i>	Have characteristic ‘double refractile wall; Usually show ‘broad based budding’ yeasts in serial/ deeper sections.
<i>Rhinosporidium Seeberi</i>	Usually show sporangia with sporangiospores in serial/deeper sections; spores stain homogeneously black in GMS; spores are ‘Carminophilic’
<i>Coccidioidomycosis</i>	Usually show sporangia with sporangiospores in serial/deeper sections; GMS stains only the wall of spores.
<i>Transverse section of Mucorales</i>	Hyphal walls of Mucorales are thin; usually show nonparallel walled right angled branching hyphae in serial/deeper sections.
<i>Helminths (Strongyloides, Diofilaria)</i>	Have thick outer cuticle; organs are seen inside.
<i>Small blood vessels</i>	Have endothelial cell lining; luminal blood cells; smooth muscle cells in the wall.

small blood vessels, and helminths such as *Strongyloides* and *Diofilaria*.

Degenerating yeasts of *B. dermatitidis* may look ‘empty’ and confused for smaller forms of ‘*adiaspores*’. Though degenerating, yeasts of *B. dermatitidis* retain the ‘double refractile’ nature of their wall. Serial and deeper sections may reveal typical tissue forms including evidence of ‘broad based budding’. When in doubt, Congo red stain is a very useful differentiating stain, where yeasts of *B. dermatitidis* are ‘Congophilic’ showing bright red stain of the wall (22). In addition, *Adiaspores* are ‘non replicating’ and show no evidence of ‘budding’. *Adiaspores*, although having a thick ‘trilaminar’ wall, are not ‘refractile’ and are congo red negative.

Spherules of both *Rhinosporidium seeberi* & *Coccidioidomycosis* show relatively thinner wall than those of *Adiaspores* and show characteristic endosporulation. Endosporulation is characteristically absent in *Adiaspiromycosis*.

Transverse sections of helminths usually show typical outer cuticle and organs inside in contrast to empty rings of *Adiaspores*.

Small blood vessels are usually confused for *Adiaspores*, but the presence of flattened endothelial cells on the inner side with dark blue nuclei and the presence of red blood cells rule out *Adiaspores* (Figure 1G) (3,14).

Transverse sections of irregular hyphae of ‘*Mucorales*’ are sometimes confused with ‘*Adiaspores*’. *Mucorales* have a thin wall in contrast to the thick wall of *Adiaspores*. *Mucorales* usually show folded large aseptate irregular non parallel hyphae in the wall of blood vessels, amidst the necrotic material and usually show right angled branching. With careful detailed examination of serial & deeper sections one may find ‘typical tissue form’. *Mucorales* and particularly

Rhizopus spp may show ‘ectosporulation’ with characteristic vesicles and released spores (conidia).

Retraction of the inner wall of *Adiaspore* may create ‘septate like partition’ (Figure 1C, 4E), simulating septating or multiplying fungal cell. With careful examination of deeper and serial sections, one may see typical non replicating ‘*Adiaspores*’.

Inner granular contents of *Adiaspores* may be confused with endospores of *rhinosporidiosis* and *coccidioidomycosis*. In ‘*Adiaspores*’, the inner granular contents are irregular heterogenous, amorphous, less numerous, may be seen abutting the inner membrane, and stain faintly gray in GMS and faintly eosinophilic to amphophilic in PAS. Endospores of *Rhinosporidiosis* and *Coccidioidomycosis* are numerous, more or less uniformly round, with a distinct wall and stain black in GMS and pink in PAS. Endospores of *Rhinosporidiosis* are mucicarmine positive while granular contents of ‘*Adiaspores*’ are mucicarmine negative (3).

CONCLUSION

Adiaspiromycosis is a rare self-limiting fungal infection and primary cutaneous involvement is a rare distinct entity. It is often an incidental finding in histopathology. Human tissue is the dead end of the life cycle for the fungi. The fungus is difficult to isolate from human clinical material because of the unique thermal dimorphism. Histopathology is the mainstay in diagnosis.

Conflict of Interest

The authors have no conflict of interest.

Authorship Contributions

Concept: SCS, Design: SCS, Data collection or processing: SCS, AS, Analysis or Interpretation: SCS, CTN, Literature search: SCS, CTN, AS, SV, Writing: SCS, Approval: SCS, SV.

REFERENCES

- Chandler FW, Kaplan W, Ajello L. Adiaspiromycosis. In: A Colour Atlas and Textbook of the Histopathology of Mycotic Diseases, London: Wolfe Medical, 1980;30-33.
- Rippon JW. Adiaspiromycosis. In: Medical Mycology 1988: The Pathogenic Fungi & Pathogenic Actinomycetes, 3rd edn, Philadelphia; WB Saunders Co, 1988;720-721.
- Pfaller MA, Diekema DJ. Unusual fungal and pseudofungal infections of humans. J Clin Microbiol. 2005;43(4):1495-504.
- Borman AM, Jiang Y, Dukik K, Sigler L, Schwartz IS, de Hoog S. Adiaspiromycosis and diseases caused by related fungi in Ajellomycetaceae. In: Seyedmousavi S, de Hoog G, Guillot J, Verweij P (eds). Emerging and Epizootic Fungal Infections in Animals, Springer International Publishing AG, 2018;147-158.
- Sigler L. Adiaspiromycosis and other infections caused by Emmonsia species. In: Hay RJ, Merz WG (eds). Topley and Wilson's microbiology and microbial infections. 10th ed. London: Arnold Hodder, 2005;809-824.
- Hughes K, Borman AM. Adiaspiromycosis in a wild European rabbit, and a review of the literature. J Vet Diagn Invest. 2018;30(4):614-618.
- Navas-Suárez PE, Sacristán C, Díaz-Delgado J, Yogui DR, Alves MH, Fuentes-Castillo D, Ospina-Pinto C, Zamana RR, Desbiez ALJ, Catão-Dias JL. Pulmonary adiaspiromycosis in armadillos killed by motor vehicle collisions in Brazil. Sci Rep. 2021;11(1):272.
- Nakano M, Yamaguchi E, Kimoto M, et al. Pathological study of adiaspiromycosis in Eurasian red squirrel (*Sciurus vulgaris orientis*) and brown rat (*Rattus norvegicus*) caught in Tokachi district, Hokkaido. Jpn J Zoo Wildl Med 2017;22:37-40.
- Matsuda K, Niki H, Yukawa A, Yanagi M, Souma K, Masuko T, Taniyama H. First detection of adiaspiromycosis in the lungs of a deer. J Vet Med Sci. 2015;77(8):981-3.
- Malatesta D, Simpson VR, Fontanesi L, Fusillo R, Marcelli M, Bongiovanni L, Romanucci M, Palmieri C, Della Salda L. First description of adiaspiromycosis in an Eurasian otter (*Lutra lutra*) in Italy. Vet Ital. 2014;50(3):199-202.
- Walsh TJ, Groll A, Hiemenz J, Fleming R, Roilides E, Anaissie E. Infections due to emerging and uncommon medically important fungal pathogens. Clin Microbiol Infect. 2004;10:48-66.
- Krückemeier S, Ramon M, Vidal E, Martino L, Burgaya J, Ribas MP, Dias-Alves A, Lobato-Bailón L, Pérez de Val B, Cabezón O, Espunyes J. Adiaspiromycoses in Wild Rodents from the Pyrenees, Northeastern Spain. J Wildl Dis. 2024;60(2):526-30.
- Buyuksirin M, Ozkaya S, Yucel N, Guldaival F, Ceylan K, Erbay Polat G. Pulmonary Adiaspiromycosis: first reported case in Turkey. Respiratory Medicine CME. 2011;4:166-9.
- Anstead GM, Sutton DA, Graybill JR. Adiaspiromycosis causing respiratory failure and a review of human infections due to *Emmonsia* and *Chryso sporium* spp. J Clin Microbiol. 2012;50(4):1346-54.
- Echavarria E, Cano EL, Restrepo A. Disseminated adiaspiromycosis in a patient with AIDS. J Med Vet Mycol. 1993;31(1):91-7.
- Stebbins WG, Krishtul A, Bottone EJ, Phelps R, Cohen S. Cutaneous adiaspiromycosis: a distinct dermatologic entity associated with *Chryso sporium* species. J Am Acad Dermatol. 2004;51(5 Suppl):S185-9.
- Kalam A, Thambiah AS. Adiaspiromycosis of human skin caused by *Emmonsia crescens*. Sabouraudia. 1979;17(4):377-81.
- Suchonwanit P, Chaiyabutr C, Vachiramon V. Primary Cutaneous *Chryso sporium* Infection following Ear Piercing: A Case Report. Case Rep Dermatol. 2015;7(2):136-40.
- Dincy PC, Meera T, Susanne PA, Promila RM. Disseminated cutaneous *chryso sporium* infection. Trop Doct. 2019;49(4):306-8.
- Jiang Y, Dukik K, Munoz JF, Sigler L, Schwartz IS, Govender NP, Kenyon C, Feng P, van den Ende BG, Stielow JB, Stchigel AM, Lu H, de Hoog S. Phylogeny, ecology and taxonomy of systemic pathogens and their relatives in Ajellomycetaceae (Onygenales): Blastomyces, Emergomyces, Emmonsia, Emmonsiiopsis. Fungal Diversity. 2018;90:245-91.
- Takehige A, Nakano M, Kondoh D, Tanaka Y, Sekiya A, Yaguchi T, Furuoka H, Toyotome T. Adiaspore development and morphological characteristics in a mouse adiaspiromycosis model. Vet Res. 2020;51(1):119.
- Axelson GK, Giorgadze T, Youngberg GA. Evaluation of the use of Congo red staining in the differential diagnosis of *Candida* vs. various other yeast-form fungal organisms. J Cutan Pathol. 2008;35(1):27-30.