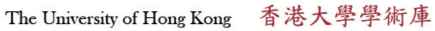
## The HKU Scholars Hub





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Citation	Journal Of Periodontal Research, 2005, v. 40 n. 3, p. 258-268
Issued Date	2005
URL	http://hdl.handle.net/10722/90726
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URL	http://hub.hku.hk/handle/10722/55446
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Characterization of Actinobacillus actinomycetemcomitans isolated from young

Chinese aggressive periodontitis patients.

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Running head: A. actinomycetemcomitans in young Chinese

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Key words: Actinobacillus actinomycetemcomitans; cdt; ltx; periodontal diseases; serotype

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Leung WK, Ngai VKS, Yau JYY, Cheung BPK, Tsang PWK, Corbet EF: Characterization of Actinobacillus actinomycetemcomitans isolated from young Chinese aggressive periodontitis patients. J Periodont Res

Objective: This study characterized Actinobacillus actinomycetemcomitans isolates from young Chinese aggressive periodontitis patients.

Methods: Subgingival plaque samples (2/subject) were collected from diseased subjects < 25 yr (n = 9,  $21.1 \pm 1.6$  yr) and age-matched periodontitis-free controls (n = 47,  $22.0 \pm 1.1$  yr). Selective and anaerobic culture were used. The serotype, leukotoxin gene (ltx) operon promoter, and the cytolethal distending toxin (cdt) genes complex of the A. actinomycetemcomitans isolates were investigated. Effects of the isolates on non-keratinzing periodontal ligament epithelial cells (PLE) monolayer were studied.

Results: Diseased subjects had significantly higher full-mouth bleeding score (P = 0.002) and TVC from plaque samples ( $7.2 \times 10^6$  vs  $2.1 \times 10^5$  c.f.u./paperpoint, P < 0.005). A. actinomycetemcomitans was isolated from 67%/56% or 6%/4% of diseased or controls subject/sites, respectively (P < 0.001). The proportion of A. actinomycetemcomitans isolatable from aggressive periodontitis or periodontitis-free associated subgingival plaque was low (0.7% vs 0.1%, P < 0.02). The serotype of the isolates was characterized. All isolates possessed 652-like ltx gene promoter and all but one serotype c isolate from a diseased patient had intact cdtABC genes. That particular strain appeared to confer the least cellular damages on PLE monolayer compared to others.

Conclusion: This preliminary study confirmed the notion of increased prevalence and quantity of A. actinomycetemcomitans associated with aggressive periodontitis in young patients. The overall ltx promoter and cdt characteristics of the A. actinomycetemcomitans isolates, however, were similar among the diseased and control groups. A strain lacking the

cdtABC gene appeared to be less damaging to a PLE cell model. Further studies, therefore, are warranted to clarify the pathogenic role and potentials of A. actinomycetemcomitans in aggressive periodontitis.

In the southern Chinese city of Hong Kong, the population incidence of deep periodontal pocket in adolescents is moderate (approx. 1%; (1, 2)) compared to 0.5-1.5% in the rest of the world ((3), review). However, the proportion of 15 – 19 year-olds in Hong Kong with shallow periodontal pockets was 26% (1), which was comparable to the 22% subjects of the same age, who had shallow pockets in the United States (4). Periodontal diseases observable in young individuals are believed to be caused by various periodontopathogens, as in the case for older individuals (5). The microorganism *Actinobacillus actinomycetemcomitans* is regarded as one of the key bacterial agents associated with aggressive periodontitis in young adults (6).

Six serotypes of A. actinomycetemcomitans, i.e. serotype a, b, c, d, e and f, have been identified (7). A. actinomycetemcomitans is a special periodontopathogen in the sense that in most strains at least two complex multigene toxin systems are in operation, the leukotoxin (RTX) (8) and the later identified cytolethal distending toxin (CDT) (9). The RTX or repeats in the structural toxins comprise a family of large, heat-labile, Ca2+-dependent, pore forming cytotoxins that display different target cell specificities (10). A. actinomycetemcomitans leukotoxin is active against lymphocytes and granulocytes (8) triggering hyper-production of reactive oxygen intermediates, and causes degranulation and release of lysosomal contents leading to inflammation-mediated injury of infected host tissue or programmed cell death in affected host cells (11). The ultimate effect of A. actinomycetemcomitans leukotoxin seems to enable the pathogen to evade the host immune system, hence establishing a local infection. CDT is a multicomponent bacterial holotoxin that targets most eukaryotic cells causing distention and cell cycle arrest (12). In brief, CdtB, the active subunit of the CDT holotoxin, through its nuclease activity, could induce limited DNA damage of infected eukaryotic cells and hence lead to arrest of cell cycle or proliferation (13). This action of CDT was suggested to be a strategy used by A. actinomycetemcomitans to moderate host cell functions (14).

In recent years, a series of clinical and microbiological studies on young Chinese adults of southern and western Chinese decent with poor oral hygiene and untreated gingivitis (15) showed a high prevalence of the putative periodontopathogens Porphyromonas gingivalis and A. actinomycetemcomitans in subgingival plaque samples (16). examination of the leukotoxin gene lktA operon of the A. actinomycetemcomitans isolates, Mombelli and co-workers (16) could not detect the presence of a deletion in the promoter region indicative of a highly leukotoxic (or JP2 type) A. actinomycetemcomitans strain. Later on, the same group conducted a larger scale microbiological study which included the above young subjects with untreated gingivitis (n = 73), 35-44 year-old subjects from an oral epidemiological survey in southern China (n = 81), and middle-aged periodontitis patients in Hong Kong (n = 31) (17). They concluded that A. actinomycetemcomitans appeared to be a common constituent of the normal oral flora in the Chinese subjects, a finding confirmed by Tan and co-workers in Singapore (18), and also shown for a rural adult population in southern Thailand (19). No highly leukotoxic strain similar to JP2 type of lktA promoter sequence could be isolated by Mombelli and co-workers in Chinese subjects (17) complementing previous reports about racial tropism of a highly leukotoxic ltx clone of A. actinomycetemcomitans (20,21) shown to be pathogenic among A. actinomycetemcomitans detected in early onset periodontitis (22).

The prevalence of CDT in A. actinomycetemcomitans clinical isolates has also recently been studied. Clinical isolates from Brazil, Kenya, Japan and Sweden were analyzed (23) and three cdt genes, i.e. cdtA, cdtB and cdtC, were detectable from 34 of 40 A. actinomycetemcomitans strains. However, that study did not report the relationship between CDT positivity and the periodontal status of the patients and the sites from which the A. actinomycetemcomitans were sampled. A later report by Tan et al. (2002) (24) using direct PCR detection of subgingival plaque samples from 146 Singaporean patients with various

periodontal diseases detected a low prevalence of the cdt genes complex. In that study, subjects of mean age  $42.5 \pm 6.5$  years were sampled. 12% of diseased sites with A. actinomycetemcomitans were cdt genes complex positive, of which 77% were from adults diagnosed with aggressive periodontitis, showing a strong association between the presence of these genes with severe disease conditions. No data, however, is available yet regarding the occurrence of A. actinomycetemcomitans cdt genes in young adults suffering from aggressive periodontitis.

The aim of the present study was hence to investigate the A. actinomycetemcomitans carrier rate in young Chinese subjects with untreated aggressive periodontitis and to compare this with the carrier rate in periodontitis-free age-matched controls, and to study the prevalence and nature of the A. actinomycetemcomitans isolates' serotypes, and the two complex multigene toxin genotypes and their pathogenic potential in a porcine periodontal ligament epithelial (PLE) cell explant model.

#### Materials and methods

#### Subjects

Subgingival microbiological samples were obtained from nine untreated aggressive periodontitis patients recruited over a period of 6 months, who also fulfilled the following inclusion criteria: i) < 25 year, ii) possessing pockets  $\ge 5$  mm on at least 2 of the first molars or incisors and these teeth should be free from > 1 surface restorations, not endodontically treated or showing signs of pulpal diseases, iii) not taking any systemic antibiotics in the preceding 6 months, and iv) not suffering from any debilitating systemic diseases and not pregnant. The aggressive periodontitis subjects were recruited from the University Health

Service Dental Clinic and the Reception Clinic of the Prince Philip Dental Hospital, Faculty of Dentistry, the University of Hong Kong. During the recruitment period, approximately 800 first year university students attended the University Health Service Dental Clinic for initial examination and six aggressive periodontitis patients satisfying the inclusion criteria were identified and recruited. Another second-year university student with aggressive periodontitis was recruited out of three potentially suitable cases referred from the University Health Service Dental Clinic for possible inclusion. Two more subjects were recruited during the same period from a total of 5,041 adult patients who attended the Reception Clinic, Faculty of Dentistry for screening. Fifty age-matched periodontitis-free university dental students (Control group) were recruited, who had i) sound (≤ 1 surface restoration, not endodontically treated or showing signs of pulpal disease) first molars and incisors, ii) no probing pocket depth > 3 mm, iii) no radiographic signs of alveolar bone loss on any standing tooth.

#### **Clinical Examination**

The following parameters were measured at six sites on all teeth: i) Plaque recording (modified from (25)), ii) probing pocket depth using William's 14W periodontal probe, and iii) bleeding on probing. Clinical attachment level was measured from the depth of the sampling pocket to the cemento-enamel junction. All aggressive periodontitis subjects, after examination and microbiological sampling, were given a comprehensive course of non-surgical periodontal therapy followed by appropriate supportive periodontal therapy. Surgical periodontal therapy was carried out for individual patients when indicated.

#### Microbiological Sampling

In the aggressive periodontitis patient group, the deepest periodontal pockets from two non-neighbouring teeth were selected for microbial sampling. All sample sites from the aggressive periodontitis group were from first molars. For control subjects, two sites were selected randomly from any two of the four first molars for microbial sampling. The microbiological sampling was performed as described previously (26). In brief, the sites to be sampled were dried and isolated with sterile cotton wool rolls. Supragingival plaque was removal from the sample site and adjacent teeth with a sterile curette. Four sterile endodontic paper points (medium absorbent points, DiaDent, Burnaby, Canada) were inserted into the sample site 2 at a time until resistance was felt and left in place for 20 seconds. The specimens were than transferred to the laboratory in 1 ml reduced transport fluid at 4°C. All samples were processed within one hour after sampling.

#### Culture

All samples were vortexed for 20 seconds at maximum setting (Autovortex Mixer SA2, Stuart Scientific, London, UK) and then serially diluted in 10-fold increments up to a thousand fold in trypticase soy broth (TSB). The diluted samples were plated onto enriched Columbia blood agar (CBABS, Columbia agar base with 5% defibrinated horse blood, 0.0005% hemin and 0.00005% Vitamin K) (26) and Tryptic Soy-Serum-Bacitracin-Vancomycin Agar (TSBV) (27) using a Spiroplater (Spiral Plater Model D, Spiral System Inc., Cinannati, OH). CBABS and TSBV plates were incubated in an anaerobic chamber (Forma Scientific Inc, Marietta, OH) under an atmosphere of 10% CO<sub>2</sub>, 10% H<sub>2</sub>, 80% N<sub>2</sub> for 5 - 7 days at 37°C.

TSBV or CBABS plates with appropriate number of colonies that were well separated and evenly dispersed with 30 – 300 colony-forming units were chosen for counting. Total viable counts (TVC) measurement was performed on CBABS plates. The presumptive

identification of A. actinomycetemcomitans on TSBV plate was based on colony morphology (transparent colonies with characteristic stellar structure) and a positive catalase reaction. Presumptive A. actinomycetemcomitans isolates were subcultured in TSBV plates to obtain pure cultures. Pure culture of A. actinomycetemcomitans was identified using MicroSeq 500 16S rDNA-based bacterial identification system (Perkin-Elmer Applied Biosystems Division, Foster City, CA). With reference to the dilution factor and the count on TSBV, the A. actinomycetemcomitans c.f.u. per paper point was also calculated.

#### **Bacterial strains**

The bacteria strains used in this study and their relevant genotypic characteristics were as shown in Table 1.

#### DNA preparation of A. actinomycetemcomitans

DNA was isolated from 2-days-old A. actinomycetemcomitans reference—strains or isolates cultured in TSB supplemented with 1% (wt/vol) yeast extract. Isolation of chromosomal DNA was carried out according to the instruction on the QIAamp DNA kit (Qiagen, Hilden, Germany). In brief, 200  $\mu$ l of the bacterial culture on TSB was pelleted at 5,000  $\times$  g for 10 min, washed once in phosphate buffered saline, PBS, pH 7.2 and then resuspended in 180  $\mu$ l ATL Buffer. Twenty microlitre of Proteinase K was added to the mixture, vortexed and incubated at 56°C on a shaking water bath overnight for cell lysis. Then 200  $\mu$ l AL Buffer was added to the sample, followed by pulse-vortexing of the mixture and incubation at 70°C for 10 min. The bacterial lysate was washed in the following sequence: i) addition of 200  $\mu$ l 99% ethanol to the sample, vortexed 15 s, then carefully applied to the QIAamp spin column, centrifuged at  $6,000 \times g$ , 1 min; ii) 500  $\mu$ l AW1 Buffer at  $6,000 \times g$ , 1 min, iii) 500  $\mu$ l AW2 Buffer at  $13,000 \times g$ , 3 min, followed by a repeated spin at  $13,000 \times g$ , 1 min. Then, 200  $\mu$ l

AE Buffer was added and the spin column incubated at room temperature for 1 min, followed by DNA elution at  $6,000 \times g$ , 1 min. The elution step was repeated by adding another 200  $\mu$ l AE Buffer and the resultant eluted DNA were pooled, aliquoted and stored at -20°C for future PCR analysis.

#### MicroSeq 500 16S rDNA-based bacterial identification

DNA extract from pure culture of presumptive A. actinomycetemcomitans strains was amplified according to the manufacturer's instruction. In brief, the 25 µl of 2 × PCR master mix was added to 25 µl of DNA template (1 µg/ml). The mixture was amplified at 95°C for 10 min initial denaturation, 30 cycles of 95°C for 30 s, 60°C for 30 s, and 72°C for 45 s, with a final extension at 72°C for 10 min in a GeneAmp 9700 machine (Applied Biosystems). The amplified product was purified using the QIAquick gel extraction kit (Qiagen). PCR products were bi-directionally sequenced with forward and reverse sequencing mix using an ABI 310 automated sequencer according to manufacturer's instruction (Applied Biosystems). The DNA sequences were analyzed using the database provided by the MicroSeq 500 16S rDNA-based bacterial identification system. The sequence homology analysis was performed using the BLAST algorithm in database of National Center for Biotechnology Information.

#### PCR analysis of A. actinomycetemcomitans genes

PCR primers

Table 2 lists the PCR primers used in this study.

Serotyping

PCR were performed on each 16S rDNA confirmed *A. actinomycetemcomitans* isolate as described by Kaplan et al (7). The primer sets used were as shown in Table 2. The 100µl volume of the PCR master mix contained 2 µl of *A. actinomycetemcomitans* DNA template, 2 µl of 50 mM MgCl<sub>2</sub>, 10 µl of 0.2 M Tris-0.5 M KCl, pH 8.4 (10× PCR buffer), a 150 µM concentration of each dNTP, 1µM of the corresponding forward and reverse primer and 2 U of Taq polymerase (Life Technologies, Frederick, Maryland). The 30 cycles of the PCR protocol included 30 s of denaturation at 94°C, 90 s of annealing at 57°C, 90 s of primer extension at 72°C. The PCR product were electrophoresed in 1.5% agarose gel in TBE buffer, stained with ethidium bromide, and visualized under UV transillumination. Isolates of the same serotype groups were noted and the analysis was repeated on at least one separate occasion with strains recovered from the stock kept at -70°C. Then the amplicon from the isolates underwent electrophoresis side-by-side with amplicon from positive *A. actinomycetemcomitans* control (strain of known serotype except for serotypes e and f which were not available) to confirm the findings.

#### Characterization of A. actinomycetemcomitans ltx operon promoter

The characteristic of the *A. actinomycetemcomitans* leukotoxin gene operon promoter was studied by PCR (20, 28) with slight modifications to their methods. The corresponding custom-synthesized primers (Gibco BRL, Hong Kong) used in the study were as shown in Table 2. A 100-μl volume of the PCR master mix contained 10 μl of *A. actinomycetemcomitans* DNA template, 10 μl of 0.5 M KCl-0.2 M Tris, pH 8.4 (10 × PCR buffer), a 150 μM concentration of each dNTP, 1.5 mM MgCl<sub>2</sub>, 1 μM of forward and reverse primer and 2 U of Taq polymerase (Life Technologies). The 30 cycles of the PCR protocol included 1 min of denaturation at 94°C, 1 min of annealing at 60°C, 2 min of primer extension at 72°C. Control tubes without template DNA or with DNA from *Haemophilus* 

aphrophilus were included in each run. The PCR products were analyzed by 0.8% agarose gel electrophoresis (20). Repeated PCR analysis was carried out for confirmation of results.

## PCR detection of cdtABC genes

The characterization of individual *cdtABC* genes from *A. actinomycetemcomitans* was carried out as described by Ahmed et al. (31). PCR was performed with 5 μl of the DNA template in 50 μl reaction mixture containing 2 mM MgCl<sub>2</sub>-50 mM KCl-10 mM Tris-HCl, pH 8.3, 2.5 U Taq polymerase (Life Technologies), 200 μM of each dNTP and 10 μM of each primer. The amplification cycle comprised 30 cycles denaturation at 94°C, 30 s, annealing at 57°C, 2 min and primer extension at 72°C, 2 min. The PCR products were visualized by gel electrophoresis on 1% agarose side-by-side with amplicon from positive controls, followed by staining using ethidium bromide and illumination with UV. The *cdtABC* genes analysis was repeated on one separate occasion with strains recovered from the stock kept at -70°C.

#### Eukaryotic cell culture

Porcine periodontal ligament epithelial (PLE) cell explants (32) were cultured in minimum essential alpha medium ( $\alpha$ MEM; Gibco BRL, Paisley, UK) with 15% fetal bovine serum (FBS, Gibco BRL) with 10,000 µg/ml, penicillin G, 10 µg/ml streptomycin, 1.2% fungizone (v/v) (Gibco BRL), 30 µg/ml amphotericin B and 25 µg/ml sodium deoxycholate in the manner described previously (33). In brief, the PLE cells were cultured on an 12-well chamber slide (Nune, Naperviller, IL, USA) until early confluency. Immediately prior to the assays, the culture medium was discarded and the monolayer washed twice with warm (37°C)  $\alpha$ MEM. The washed PLE cultures were challenged with 100 µg/ml of 2.5 × 10<sup>9</sup>, 5 × 10<sup>8</sup> and 1 × 10<sup>8</sup> A. actinomycetemcomitans per ml suspensions in 15% FBS –  $\alpha$ MEM, 37°C, 5% CO<sub>2</sub> in air, 24 hrs. PLE passages 5 to 10 were used in the experiment.

## Morphometric analysis of A. actinomycetemcomitans-induced PLE cell damage

A. actinomycetemcomitans-treated PLE cells were fixed and stained with May-Grünwald and Giemsa stains (E. Merck, Darmstadt, Germany) as described earlier (34). After mounting, the specimens were examined under a microscope at 40 times magnification fitted with a digital camera (Leica DC 300 V 2.0, Leica, Wetzlar, Germany). The morphometric analysis was done as previously described (33) using an image analyzing software (Qwin version 2.4, Leica, Cambridege, UK). Cell number, cell area and mean cell size were measured.

#### **Statistics**

Data on patient demography, clinical conditions, A. actinomycetemcomitans isolation, and A. actinomycetemcomitans-induced PLE cell damage - in terms of confluency, cell size, proportion of cells left attached - were expressed as mean  $\pm$  standard deviation. Differences between the means were analyzed by unpaired t-test with Welch correction, or analysis of variance with Bonferroni adjustment for multiple comparison when appropriate. Fisher's exact test was also used. Groups were regarded as significantly different from each other if P < 0.05.

#### Results

#### Clinical findings

Nine aggressive periodontitis patients fulfilled the inclusion criteria and were recruited. Forty-seven out of 50, pre-screened to meet the inclusion criteria, university dental students participated as controls. Two out of 9 and 20 out of 47 of the diseased and periodontitis-free

controls respectively were male. The age of the test (19 - 24 yr) and control (20 - 24 yr) subjects was well matched. On the whole, the diseased and periodontitis-free control subjects appeared to have similar plaque levels: 16 - 52% in aggressive periodontitis group vs 7 - 60% in healthy controls. The aggressive periodontitis subjects, however, showed significantly higher BOP%, 23 - 97%, compared to 9 - 76% in healthy controls (Table 3), indicating that periodontal or gingival inflammation was generalized among the diseased subjects. The diseased subjects possessed a mean of 7 pockets  $\geq 6$  mm (2 - 17 pockets per subject). The periodontal conditions of the 112 sample sites were as shown in Table 4. All except one site in diseased group bled on probing while only 40 sample sites (43%) from control group bled on probing.

#### Microbiological findings

Significantly higher TVC/paper point was detected in aggressive periodontitis samples than in periodontitis-free controls and the specific counts for A. actinomycetemcomitans isolates were higher in aggressive periodontitis patients than in controls (Table 4). Per subject and per site A. actinomycetemcomitans isolation prevalence was significantly higher in aggressive periodontitis subjects than in controls. Overall, the percentage proportion isolation of A. actinomycetemcomitans detected by culture was at a level below 1%.

A total of 14 A. actinomycetemcomitans strains were identified and characterized (Table 5). When more than one A. actinomycetemcomitans isolates were recovered from a subject, the genotypic profile of the isolates appeared identical. Isolates of serotype a, c and f were isolated from aggressive periodontitis sites while serotypes b and c were isolated from subgingival plaque of controls (Table 5, Fig. 1). Serotype c, which was isolated in 5 of 9 A. actinomycetemcomitans positive subjects or 8 of 14 A. actinomycetemcomitans positive sites, was the predominant serotype isolated.

All A. actinomycetemcomitans isolates possessed the 652-like Itx promoter (Table 5, Fig. 2). Only from one strain of an aggressive periodontitis patient was the cdt genes complex not detectable (Table 5, Fig. 3).

## A. actinomycetemcomitans induced PLE cell monolayer changes

A. actinomycetemcomitans reference strains and isolates induced chages in PLE cell monolayers (Table 6). When used in concentrations of 2.5 × 10<sup>9</sup> or 5 × 10<sup>8</sup> bacteria/ml, all specimens showed close to or 100% detachment of PLE cells (data not shown). At concentration of 1 × 10<sup>8</sup> bacteria/ml, all A. actinomycetemcomitans strains caused reduction in cell remaining attached, general increase in cell size and various levels of change in PLE cell confluency. A. actinomycetemcomitans strain HK 928 which possessing cdtABC gene and the JP2 ltx promoter (Table 1) appeared to induce the most damage to the PLE monolayer culture, while isolate P9 possessing 652-like ltx promoter and lacking the cdtABC gene caused least damage to the monolayer (Table 6).

#### Discussion

A. actinomycetemcomitans prevalence from subgingival plaque samples reported in earlier surveys of young subject groups, who were of comparable age to the present study's groups, ranged from around 10% for Europeans (35, 36) to more than 30% in Asians (16, 37). These studies, however, did not provide breakdown of the A. actinomycetemcomitans prevalence data in subgingival plaque with respect to clinical conditions, i.e. periodontal health, presence or absence of shallow, moderate or deep pockets, at sample sites. A German study of young adults with minimal periodontal disease, in which each tooth was sampled, found A.

actinomycetemcomitans to be associated with some deviation from gingival health (36). An Italian study reported A. actinomycetemcomitans subject/site prevalence of 4.2%/1.0% from subgingival plaque of young periodontally healthy individuals before commencement of orthodontics (38). The background of the subjects was similar to those of the controls of this report, i.e. individuals with good to fair oral hygiene without periodontitis. Both the Italian study and the current study showed similar low levels of A. actinomycetemcomitans prevalence in young individuals free from periodontitis. The university dental students who formed the control group of the present study were not all in their clinical years of study and their plaque control levels were not optimal, but their general bleeding on probing levels were lower than those found in the aggressive periodontitis patients. In young adults with A. actinomycetemcomitans it has been suggested that a heavy load of A. actinomycetemcomitans might suppress gingival inflammation (36), an effect not evident in the present study. An eight year longitudinal study from Norway followed up the periodontal status of 206 14-yearolds until they were 22 years-old. A total of 13 were found to have experienced periodontal bone loss at the end of the study. Thirteen periodontally healthy individuals were then randomly selected to be the control subjects and one (7.7%) of them carried subgingival A. actinomycetemcomitans detectable at their last review appointment (39). The reported A. actinomycetemcomitans prevalence from control subjects was at a similar level to the prevalence in periodontitis-free controls in the current study.

The findings regarding subgingival A. actinomycetemcomitans prevalence in young adults with aggressive periodontitis of the present report are similar to those reported by other studies (39,40). The mean proportion of the A. actinomycetemcomitans isolated from aggressive periodontitis associated plaque in the present study, however, appeared to be slightly lower than what has been reported previously (40). The study subjects themselves displayed the clinical conditions compatible with a classification of aggressive periodontitis

(41), although no enquiry was made about a family history of periodontitis to confirm any familial aggregation.

It has been reported that serotypes a, b, c, and f occur more frequently among oral isolates than serotypes d and e (29, 42). Serotype c seems to be associated more often with clinically healthy situations, although A. actinomycetemcomitans of serotypes a, b, c, f could all be detected from diseased subgingival plaque specimens (7), suggesting that phylogenetically diverse strains carry pathogenic potential (43). Serotypes a, b, c and f of A. actinomycetemcomitans strains were isolated from both periodontitis-free control and aggressive periodontitis plaque samples in the present study (Table 5), reconfirming what was observed earlier (29, 42). For Japanese adults it was suggested that serotype c, which was the most common serotype found in the periodontitis-free controls, may differ from other serotypes (44). In one out of eight serotype c A. actinomycetemcomitans isolates, one strain from a diseased patient lacked the cdtABC gene and was found to be less damaging to the PLE monolayer (Tables 5 & 6). The other serotype c isolates, be it from healthy or disease patients, however, possess similar pathogenic properties in PLE monolayer. Further investigation, therefore, is needed to clarify the role of serotype c in periodontal health and disease.

As was found previously with respect to A. actinomycetemcomitans isolated from Chinese subjects of different ages and clinical status (16,18), a highly leukotoxic genotype, considered to be an specific association with aggressive periodontitis in many populations (45) was not found in the present study, further confirming the belief that this highly leukotoxic genotype may be population specific (17, 21, 22). The per subject prevalence of 652 ltx promoter positive A. actinomycetemcomitans in subgingival plaque of Chinese migrant workers (16) and Singaporean Chinese adults (18), however, is very much different than reported in the current study. We postulate that the high plaque level (<25% tooth-site plaque

free) of the migrant Chinese workers (15) might reflect less healthy gum condition and hence higher chances for A. actinomycetemcomitans detection (35, 36), while the discrepancies between the A. actinomycetemcomitans prevalence in current control group than in the older healthy Singaporean Chinese remain to be elucidated.

The high prevalence current study shows of CDT genotype actinomycetemcomitans from plaque in aggressive periodontitis or periodontitis-free young individuals (Table 5). This contrasts with an earlier report showing lower per site subgingival plaque PCR detection rate (30% versus 50% in the present study) of the A. actinomycetemcomitans CDT genotype from generalized early-onset and refractory periodontitis patients, mean age 43 years, in Singapore (24). The small number of young aggressive periodontitis subjects in the present study, however, precludes a definitive explanation for these different observations. The small number of aggressive periodontitis subjects recruited resulted from the difficulties in detecting untreated cases who met all the inclusion criteria listed. However, these were young patients with a mean age of 21 years. compared to the patients of Tan and co-workers (24) who had a mean age of 43 years. A recent report studying A. actinomycetemcomitans in an older Hong Kong Chinese periodontitis patient cohort showed comparable per subject but lower per site (38% vs 56%) A. actinomycetemcomitans prevalence (46). These considerations might help in explaining why there was a lower per site A. actinomycetemcomitans CDT gene complex detection by Tan and coworkers (24) than in the current study.

To elucidate the exact relation of the different A. actinomycetemcomitans genotypes isolated in association with periodontal health and disease, one needs to study the in vivo expression of the two virulent genes studied, i.e. Itx and cdtABC which was not done in the present study. Rather, the in vitro effects of A. actinomycetemcomitans whole cell were studied. The PLE cell monolayer model enabled in vitro evaluation of pathogenic potentials

of the *A. actinomycetemcomitans* isolates. The extent of PLE cells damage appeared to correlate somewhat with the characterics of the *ltx* operon promoter and *cdtABC* genes complex (Tables 5 and 6). Caution must be exercised in interpretation of the preliminary cytopathic data of this present study. Recently reported pathogenic mechanisms of *A. actinomycetemcomitans* were not investigated. The adherence functions of the *tad* gene locus (47) or *Aae* gene (48), the apoptotic effector *cagE* homologue (49), T-cell apoptotic GroEL-like molecules (50), and other yet unknown virulent genes or factors of the periodontopathogen, could also contribute to the whole bacterium induced *in vitro* cell damage, which were not characterized in the present study.

It is well known that periodontitis develops when a susceptible host is challenged by periodontopathogens (5). It is likely that, a consortium of bacteria participate in the induction of both chronic and aggressive periodontitis. The current investigation studied the factors of a single peridontopathogen in isolation, without considering the contributions of any other periodontopathogens which might co-exist in the clinical situation. Obviously, colonization patterns by A. actinomycetemcomitans and characterization of their ltx and cdtABC genes alone are not enough to explain differences in possible pathogenic potentials of this microbe. Therefore, the results of the current study can only explain part of contribution of only one pathogen in aggressive periodonitits. Furthermore, the influence of a susceptible host towards the pathogen-human interaction was not evaluated in this current study. The reason why one cdtABC deficient A. actinomycetemcomitans strain was isolated from an aggressive periodontitis individual while the other periodontitis-free or disease associated strains with intact cdtABC gene could perhaps be explained by the above arguments. Nevertheless, the present study did provide preliminary evidence showing variations of in vitro pathogenic potentials related to the virulent genes of the periodontopathogen A. two actinomycetemcomitans.

The present preliminary study has shown an elevated prevalence and quantity of the periodontopathogen A. actinomycetemcomitans detected in subgingival plaque using selective culture in young untreated aggressive periodontitis individuals. This finding indicates that the bacterium is associated with the clinical situation and further confirms the common belief that it plays a part in the aggressive periodontitis disease process. The investigated three genotypic characteristics among the A. actinomycetemcomitans isolates, however, did not pin-point an association of any particular A. actinomycetemcomitans genotype studied with periodontitis-free status or aggressive periodontitis. The present study focused on two widely studied virulent genes of this periodontopathogen, while the genotypes in relation to adherence abilities (47, 48) or host cell apoptosis induction factors (49, 50) of the isolates were not studied. The in vitro effects of the whole-cell A. actinomycetemcomitans isolates on PLE monolayer in the present study appeared to correspond to the characteristics of these two virulent genes studied. However, the possibility still exists of variations in carriage of other virulent genes in A. actinomycetemcomitans isolates from different clinical situations, which might underpin differences in pathogenic potential of this pathogen. To fully elucidate the pathogenic role in periodontitis of the two A. actinomycetemcomitans virulent genes, including the two investigated in the present report, study of the control and/or in vivo expression of the genes in subgingival plaque associated with different clinical states is required. Laboratory study of various purified virulent genes' products and their in vitro actions are often required for understanding exact pathogenic mechanisms. Therefore, further studies are needed before the pathogenic potentials of A. actinomycetemcomitans in various periodontal disease states could be further clarified.

## Acknowledgements

We thank Dr. Mogens Kilian, University of Aarhus, Denmark for providing A. actinomycetemcomitans and H. aphrophilus strains. We also wish to thank Dr. Chun Hung Chu, University Health Service, Dental Clinic, The University of Hong Kong, for assistance in subject recruitment and Mr. Y.Y. Chui for technical assistance in the PLE experiments. This study was supported by the University of Hong Kong Committee on Research and Conference Grants No. 10201254 and 10201296.

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Table 1. Bacterial strains.

	ltx promoter Serotype class <sup>a</sup>	cdt genotype <sup>b</sup>			AND	
Strains		class <sup>a</sup>	$\overline{cdtA}$	cdtB	cdtC	Source
Actinobacillus acti	inomycetemco	omitans	***************************************	······································		
ATCC 29522	b	652	+	+		American
ATCC 29523	a	652	+	+	+	Туре
ATCC 33384	С	652	+	+	+	Culture
ATCC 43718	b	652	+	-	+	J Collection
JP2	ь	JP	+ .	+	+	_
HK 928	d	JP		+	+	Dr. Mogens
HK 929	ь	JP	+	+	+	Kilian,
HK 1037	b	652	+	+	+	University of Aarhus
Haemophilus aphr	ophilus					Aanus
HK 329						J

<sup>&</sup>lt;sup>a</sup> Promoters were classified as 652-like or JP2-like on the basis of PCR amplification of the promoter region as described by Brogan et al. (28); promoter class was named as published previously (28,29) and/or as detected by the current study.

<sup>6</sup> Genotype as described by Yamano et al. (30) and/or as detected by the current study.

Table 2. PCR primers used in characterization of Actinobacillus actinomycetemcomitans.

Primer	Sequence (5' - 3')	Analysis	Product (base)	Reference
I. Serotype				•
P11	TCTCCACCATTTTTGAGTGG	Serotype b	333	(7)
P12	GAAACCACTTCTATTTCTCC	Serotype c	268	<b>X</b> /
P13	CCTTTATCAATCCAGACAGC	Serotype f	232	
P14	ARAAYTTYTCWTCGGGAATG <sup>a</sup>	w <b>3</b>		
P15	TGGGTCATGGGAGGTACTCC	Serotype a	293	
P16	GCTAGGAACAAAGCAGCATC	× #.		
<sup>6</sup> P17	TGGAACGGGTATGGGAACGG	Serotype d	411	
P18	GGATGCTCATCTAGCCATGC			
P19	ATTCCAGCCTTTTGGTTCTC	Serotype e	311	
P20	TGGTCTGCGTTGTAGGTTGG	- January P	~	
II. Leukotoxin gen	ne operon promoter			
Forward	TTTCTCCATATTAAATCTCCTTGT	ltx promoter	504 or 1034	(20)
Reverse	CAGATCAAAACCTGATAACAGTATT		00,01,00	(20)
III. Cytolethal dist	ending toxin			
cdtA-7	GATGGATCTAAGGAGAGATATAATG	cdtA	326	(31)
cdtA-13	AATTAACCGCTGTTGCTTCTAATACAG		<del></del>	(0.1)
cdtA-12	AAGGAGTTTATATGCAATGGGTAAAG	cdtB	462	
cdtA-8	TAGCGATCACGAACAAAACTAACAG			
cdtA-1	TAGTTTTGTTCGTGATCGCTAAGGAG	cdtC	272	
cdtA-4	GCTACCCTGATTTCTTCGCACCG			

 $<sup>^{</sup>a}$  R = A or G; Y = C or T; W = A or T.

Table 3. Demographic and periodontal conditions of subjects.

	Aggressive Periodontitis	Control
n	9	47
% male	22	43
Age (year, mean $\pm$ S.D.)	$21.1 \pm 1.6$	22.0 ± 1.1
Plaque % (mean ± S.D.)	$27.8 \pm 15.6$	26.4 ± 14.7
BOP % (mean $\pm$ S.D.) <sup>a</sup>	$56.8 \pm 25.9$	$28.9 \pm 22.5$
No. of pocket $\geq 6$ mm (mean $\pm$ S.D.)	$6.8 \pm 4.9$	0

<sup>&</sup>lt;sup>a</sup> Significant difference between aggressive periodontitis subjects and control group, unpaired t-test, P = 0.0016.

Table 4. Clinical and microbiological data

	Aggressive	
	Periodontitis	Control
Sample site	·	
BOP %a	94%	43%
Pocket depth $(mean \pm S.D.)^b$	$6.7 \pm 1.7$	$1.9 \pm 0.6$
Clinical attachment level (mean $\pm$ S.D.) <sup>b</sup>	$6.8 \pm 2.0$	$1.9 \pm 0.6$
Total anaerobic viable count (× 10 <sup>6</sup>		
c.f.u./paper point, mean ± S.D.) <sup>b</sup>	$7.2 \pm 9.06$	$0.2 \pm 0.5$
Actinobacillus actinomycetemcomitans		
Prevalence per subject <sup>a</sup>	67%	6%
Prevalence per site <sup>a</sup>	56%	4%
No. of strains per subject <sup>b</sup>	$1.0 \pm 0.9$	$0.1 \pm 0.4$
Count (c.f.u./paper point) <sup>b</sup>	$2.1 \times 10^4 \pm 3.1 \times 10^4$	$6.3 \times 10^2 \pm 3.7 \times 10^3$
Percentage proportion <sup>b</sup>	$0.7 \pm 0.6$	$0.1 \pm 0.8$

 $<sup>^{\</sup>rm a}$  Significant difference between aggressive periodontitis and periodontitis-free groups, P <

<sup>0.0002,</sup> Fisher's exact test. 
b Significant difference between aggressive periodontitis and periodontitis-free groups, P < 0.02, unpaired t-test, with Welch correction.

Table 5. Genotypic characteristics of Actinobacillus actinomycetemcomitans isolates<sup>a</sup>.

	· · · · · · · · · · · · · · · · · · ·		Cdt genotype <sup>d</sup>	
Sample site	Serotype <sup>c</sup>	A	В	C
periodontitis				
16ML	c	+	+	+
46DB	c	where	. +	+
26MB	c	+	+	. +
36DB	c	+		+
26ML	a	+	a few	+
46ML	a	+	w <del>y</del> w	+
16MB	f	+	+	+
26MB	f	+	+	+
46MB	f	-	+	+
36MB	c	- Oper	•	one:
16DL	c	+	+	+
46MB	b	+	nĝo.	*
26DB	c	+	+	+
46DL	c	<del></del>	» figur	+
	periodontitis 16ML 46DB 26MB 36DB 26ML 46ML 16MB 26MB 46MB 36MB	Deriodontitis         16ML       c         46DB       c         26MB       c         36DB       c         26ML       a         46ML       a         16MB       f         26MB       f         46MB       f         36MB       c         16DL       c         46MB       b         26DB       c	Deriodontitis  16ML	Sample site         Serotype <sup>c</sup> A         B           Deriodontitis         16ML         c         +         +         +         46DB         c         +         +         +         +         +         46DB         c         +

<sup>&</sup>lt;sup>a</sup> Identity of all *A. actinomycetemcomitans* isolates were confirmed by 16S rDNA partial sequencing (MicroSeq 500, Applied Biosystems) and all of them process the 652-like *ltx* promoter (20).

<sup>&</sup>lt;sup>b</sup> A. actinomycetemcomitans strains designation: P = isolates from aggressive periodontitis patients, P-F = isolates from periodontitis-free controls; the Arabic numeral following the letters P or P-F denotes the code for an individual subject; the small letter a or b denotes isolates from different sample sites of the same subject.

<sup>&</sup>lt;sup>c</sup> Serotype detection as per Kaplan et al. (7).

<sup>&</sup>lt;sup>d</sup> += the *cdt* A, B, or C gene detectable by PCR; -= the *cdt* A, B, or C gene not detectable by PCR (31).

Table 6. Effect of Actinobacillus actinomycetemcomitans isolates on porcine periodontal ligament epithelial (PLE) cell monolayer cultures<sup>a</sup>.

	PLE°				
strains <sup>b</sup>	Cell size (µm²)	Proportion of cells remaining attached (%) <sup>d</sup>	Culture confluency (%) <sup>e</sup>		
Untreated monolayer	1974 ± 1629	100 ± 13.5	98 <u>+</u> 4.6		
Aggressive periodontitis					
Pla/b	$3638 \pm 3181$	$55.7 \pm 15.1$	$96.3 \pm 4.8$		
P4a/b	2746 ± 1995	67.8 ± 19.6	$98.7 \pm 3.5$		
P5a/b	5146 ± 3268	$40.9 \pm 12.9$	85.8 ± 16.6		
P7a/b	$4133 \pm 3174$	$47.5 \pm 12.2$	93.8 ± 2.9		
P8	$4001 \pm 3252$	$51.0 \pm 15.0$	94.5 ± 5.8		
<b>P</b> 9	$2436 \pm 1784$	$78.4 \pm 14.9$	97.5 ± 8.8		
Controls					
P-F12	2864 ± 1999	46.6 ± 14.7	$92.3 \pm 9.3$		
P-F33	3518 ± 3009	54.1 <u>+</u> 14.5	$96.1 \pm 12.4$		
P-F45a/b	3103 ± 2185	53.1 ± 8.1	$94.8 \pm 6.7$		
Reference strains					
ATCC 29522	3716 ± 3197	52.8 ± 17.5	94.7 ± 13.9		
ATCC 29523	5430 ± 4209	36.5 + 13.6	77.5 + 12.9		
ATCC 33384	3915 ± 3239	53.1 ± 9.9	$93.3 \pm 10.8$		
HK 928	$2158 \pm 1536$	$18.5 \pm 5.3$	$23.9 \pm 6.2$		

\*PLE monolayers were treated with  $1 \times 10^8$  cells/ml in 15% FBS- $\alpha$ MEM for 24h

<sup>&</sup>lt;sup>b</sup>A. actinomycetemcomitans strains designation: please refer to Table 5 for details. Isolate a or b from the same patient, i.e. P1, P4, P5, P7 or P-F45 were found to possess identical serotype, ltx promoter, cdt genotype and similar effects on PLE monolayers and their data were pooled.

<sup>°</sup>PLE stained with May-Grünwald and Giemsa stains (34); shown are mean  $\pm$  SD; n = 6 or 12 (pooled data) obtained from two independent experiments.

dSignificantly different (P < 0.05, Bonferroni multiple comparison) data values between: untreated monolayer vs all except strain P9 treated PLE; P9 treated PLE vs P5a/b, P7a/b, P-F12, ATCC 29523 and HK 928 treated PLE; HK 928 treated PLE vs all except P5a/b, P-F12 and ATCC 29523 treated PLE.

<sup>&</sup>lt;sup>e</sup>Significantly different ( $P \le 0.05$ , Bonferroni multiple comparison) data values between HK 928 treated PLE vs the rest.

### Legend

Fig. 1. PCR determination of A. actinomycetemcomitans isolates serotype. The primers listed in Table 2 were used (7). Amplicons were visualized by gel electrophoresis in 1.0% agarose. Shown are examples from different A. actinomycetemcomitans strains/isolates. Lane M1: DNA ladder molecular weight standard (MBI Fermentus); lanes 1 and 2, A. actinomycetemcomitans serotype a: ATCC 29523 and strain P5a; lanes 3, and 4, serotype b: ATCC 29522 and strain P-F33; lanes 5 and 6, serotype c: ATCC 33384 and strain P1a; lane 7, serotype d: HK 928; lane 8, serotype f: strain P7a; lane M2: 1 kb DNA ladder molecular weight standard (MBI Fermentus). Note that i) PCR amplicon sizes for various serotypes: a = 293 base-pairs, b = 333 base-pairs, c = 268 base-pairs, d = 411 base-pairs, f = 232 base-pairs; ii) no serotype d nor e was detected from the isolates recovered; iii) no reference strain profiles for serotype e or f is shown.

Fig. 2. PCR determination of the leukotoxin gene (Itx) promoter size of A. actinomycetemcomitans isolates. The primers listed in Table 2 were used (20). Amplicons were visualized by gel electrophoresis in 0.8% agarose. Shown are examples from A. actinomycetemcomitans strains/isolates carrying 652-like (1034 base-pairs, lanes 2 – 5, 9 –11) or JP2-like (504 base-pairs, lanes 6 – 8) promoters. A. actinomycetemcomitans strains isolated in the current study all carried 652-like promoter. Lane 1: DNA ladder molecular weight standard (MBI Fermentus); lanes 2 – 11: A. actinomycetemcomitans strains P-F33, P1a, P5a, P7a, JP2, HK 928, HK 929, ATCC 29522, ATCC 29523, and ATCC 33384 respectively; lane 12: H. aphrophilus HK 329; lane 13: milli Q water control.

Fig. 3. Detection of cytolethal distending toxin (cdt) genes complex of A. actinomycetemcomitans isolates by PCR. The primers listed in Table 2 were used (31). Amplicons were visualized by gel electrophoresis in 1.0% agarose. Shown are examples from different A. actinomycetemcomitans strains/isolates. Lane M1: DNA ladder molecular weight standard (MBI Fermentus); lanes 1: A. actinomycetemcomitans strain P-F33; lanes 2: strain P1a; lanes 3: strain P7a; lanes 4: strain P9; lanes 5: ATCC 29523; lane M2: 1 kb DNA ladder molecular weight standard (MBI Fermentus). Note that no cdtABC genes complex is detectable from isolate P9 while the remaining A. actinomycetemcomitans strains all showed typical sized PCR products, i.e. cdtA = 326 base-pairs, cdtB = 462 base-pairs, and cdtC = 272 base-pairs.

